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(54) Title: METHODS AND COMPOSITIONS FOR TREATING INTERVERTEBRAL DISC DEGENERATION

## Histology- Pilot Study #1



Untreated disc



Control



Treated disc

## 2 Months Post-Injection

- Untreated disc exhibits extensive degeneration
- Cross-linked matrix/BP-treated disc retains

normal structures similar to Control disc

(57) Abstract: A fluid matrix comprising cross-linked remodelable collagen from a donor vertebrate animal is useful for regenerating hydrodynamic function in damaged intervertebral discs in vivo. The matrix may be injectable and may comprise cells and a plurality of purified cell growth factors. The matrix promotes cell growth and elaboration of proteoglycans to facilitate regeneration of native tissues. The collagen in the matrix may be cross-linked using photooxidative catalysis and visible light, and purified cell growth factors are preferably at least partly bone-derived:

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## Methods And Compositions For Treating Intervertebral Disc Degeneration

### Description

#### 5 Background Art

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This invention relates generally to methods and compositions useful in treating intervertebral disc impairment in humans and other mammals. More particularly, this invention concerns compositions useful in restoring hydrodynamic function and stimulating cell proliferation and extracellular matrix production in intervertebral discs that have been compromised by injury, degenerative disease, congenital abnormalities, and/or the aging process.

Compositions of the invention may be injectable, and may include growth factors, bioactive agents, and living cells. The compositions are useful for restoring, improving, or augmenting hydrodynamic function of the intervertebral disc, increasing intervertebral disc height, and stimulating cell proliferation and/or extracellular matrix production in intervertebral discs.

The human vertebral column (spine) comprises a plurality of articulating bony elements (vertebrae) separated by soft tissue intervertebral discs. The intervertebral discs are flexible joints which provide for flexion, extension, and rotation of the vertebrae relative to one another, thus contributing to the stability and mobility of the spine within the axial skeleton.

The intervertebral disc is comprised of a central, inner portion of soft, amorphous mucoid material, the nucleus pulposus, which is peripherally surrounded by an annular ring of layers of tough, fibrous material known as the annulus fibrosus. The nucleus pulposus and the annulus fibrosus together are bounded on their upper and lower ends (i.e., cranially and caudally) by vertebral end plates located at the lower and upper ends of adjacent vertebrae. These end plates, which are composed of a thin layer of hyaline cartilage, are directly connected at their peripheries to the lamellae of the inner portions of the annulus fibrosus. The lamellae of the outer portions of the annulus fibrosus connect directly to the bone at the outer edges of the adjacent vertebrae.

The soft, mucoid nucleus pulposus contains chondrocytes, which produce fibrils of collagen (primarily Type II collagen, but also Types IX, XI, and others) and large molecules of negatively charged, sulfated proteoglycans, as depicted in Figure 1. The term matrix as used herein refers to a composition which provides structural support for, and which facilitates respiration and movement of nutrients and water to and from, an intervertebral disc. The collagenous components of the nucleus pulposus extracellular matrix comprise a scaffold that provides for normal cell (i.e., chondrocyte) attachment and cell proliferation. The negatively

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charged proteoglycan component of the nucleus pulposus extracellular matrix attracts water to form a hydrated gel, which envelops the collagen fibrils and chondrocyte cells. In the normal healthy nucleus pulposus, water comprises between 80-90% of the total weight.

The nucleus pulposus thus plays a central role in maintaining normal disc hydrodynamic function. The large molecular weight proteoglycans are contained within the nucleus pulposus by the annulus fibrosus and by the vertebral end plates, and they attract water into the nucleus through sieve-like pores in the end plates. The resulting osmotic pressure within each disc tends to expand it axially (i.e., vertically), driving the adjacent vertebrae further apart. On the other hand, mechanical movements resulting in axial compression, flexion, and rotation of the vertebrae exert forces on the intervertebral discs, which tends to drive water out of the nucleus pulposus. Water movements into and out of an intervertebral disc under the combined influence of osmotic gradients and mechanical forces constitute hydrodynamic functions important for maintaining disc health.

Movement of solutes in the water passing between discs and vertebrae during normal hydrodynamic function facilitates chondrocyte respiration and nutrition within the discs. This function is critical to chondrocyte survival since nucleus pulposus tissues of intervertebral discs are avascular (the largest such avascular structures in the human body). Maintaining sufficient water content in the nucleus pulposus is also important for absorbing high mechanical (shock) loads, for resisting herniation of nucleus pulposus matter under such loads, and for hydrating the annulus fibrosus to maintain the flexibility and strength needed for spine stability.

Normal hydrodynamic functions are compromised in degenerative disc disease (DDD). DDD involves deterioration in the structure and function of one or more intervertebral discs and is commonly associated with aging and spinal trauma. Although the etiology of DDD is not well understood, one consistent alteration seen in degenerative discs is an overall decrease in proteoglycan content within the nucleus pulposus and the annulus fibrosus. The loss in proteoglycan content results in a concomitant loss of disc water content. Reduced hydration of disc structures may weaken the annulus fibrosus, predisposing the disc to herniation. Herniation frequently results in extruded nucleus pulposus material impinging on the spinal cord or nerves, causing pain, weakness, and in some cases permanent disability.

Because adequate disc hydration is important for stability and normal mobility of the spine, effective treatment of DDD would ideally restore the disc's natural self-sustaining hydrodynamic function. Such disc regeneration therapy may require substantial restoration of cellular proteoglycan synthesis within the disc to maintain the hydrated extracellular matrix in the nucleus pulposus. Improved hydrodynamic function in such a regenerated disc may result in

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restoration and reestablishment of intervertebral disc height. It may also provide for improved hydration of the annulus fibrosus, making subsequent herniation less likely.

Prior art approaches to intervertebral disc problems fail to restore normal self-sustaining hydrodynamic function, and thus may not restore normal spinal stability and/or mobility under high loads. One approach to reforming intervertebral discs using a combination of intervertebral disc cells and a bioactive, biodegradable substrate is described in U.S. patent number 5,964,807 to Gan et al., incorporated herein by reference. The biodegradable substrate disclosed in Gan et al., including bioactive glass, polymer foam, and polymer foam coated with sol gel bioactive material, is intended to enhance cell function, cell growth and cell differentiation. The bioactive glass contains oxides of silicon, sodium, calcium and phosphorus. The polymer foam is described as biocompatible and includes polyglycolide (PGA), poly (D,L-lactide) (D,L-PLA), poly(L-lactide) (L-PLA), poly(D,L-lactide-co-glycolide) (D,L-PLGA), poly(L-lactide-coglycolide) (L-PLGA), polycaprolactone (PCL), polydioxanone, polyesteramides, copolyoxalates, and polycarbonates. Gan et al. describes application of this approach to intervertebral disc reformation in mature New Zealand rabbits, concluding with ingrowth of cells and concurrent degradation of implanted material with little or no inflammation. However, degradation of portions of the implanted material, such as acidic breakdown of PLAs, PGAs and PLGAs, may adversely affect cell growth, cell function and/or cell differentiation.

A somewhat analogous disclosure relating to tissues for grafting describes matrix particulates comprising growth factors that may be seeded with cells; see U.S. patent number 5,800,537 to Bell, incorporated herein by reference. The matrix and cells are applied to scaffolds, which include biodegradable polymers, microparticulates, and collagen which has been cross-linked by exposure to ultraviolet radiation and formed to produce solids of foam, thread, fabric or film. The matrix particulates are derived from tissue from which cells and cell remnants have been removed without removing factors necessary for cell growth, morphogenesis and differentiation. Bell specifically avoids the use of reagents like high salt, or deliysidation reagents such as butanol/ether or detergents. Such reagents are unfavorably characterized as being responsible for removing from the source tissue factors essential for stimulating repair and remodeling processes. Alternative approaches, in which such factors are obtained from other sources rather than being retained in the tissue, are not addressed.

Still another disclosure related to regeneration of cartilage is found in U.S. patent number 5,837,235 to Mueller et al., incorporated herein by reference. Mueller et al. describes comminuting small particles of autologous omentum or other fatty tissue for use as a carrier, and adding to the carrier growth factors such as Transforming Growth Factor Beta and Bone

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Morphogenic Protein. Mueller et al. does not teach cross-linking tissues to create a cross-linked matrix.

The Gan et al. patent above is representative of past attempts to restore or regenerate substantially natural hydrodynamic disc function to intervertebral discs, but such techniques have not been proven in clinical trials. Similarly, the approaches of Bell and Mueller et al. have not been widely adapted for disc regeneration, and better approaches are still needed because low back pain sufficient to prevent the patient from working is said to affect 60% to 85% of all people at some time in their life. In the absence of safer and more efficacious treatment, an estimated 700,000 discectomies and 550,000 spinal fusions are performed worldwide each year to treat these conditions. Several prosthetic devices and compositions employing synthetic components have also been proposed for replacement of degenerated discs or portions thereof. See, for example, U.S. patent numbers 4,772,287, 4,904,260, 5,047,055, 5,171,280, 5,171,281, 5,192,326, 5,458,643, 5,514,180, 5,534,028, 5,645,597, 5,674,295, 5,800,549, 5,824,093, 5,922,028, 5,976,186, and 6,022,376.

A portion of the disc prostheses referenced above comprise hydrogels which are intended to facilitate hydrodynamic function similar in some respects to that of healthy natural discs. See, for example, U.S. patent Number 6,022,376 (Assell et al.). These prosthetic hydrogels, however, are not renewed through cellular activity within the discs. Thus, any improvement in disc hydrodynamic function would not be self-sustaining and would decline over time with degradation of the prosthetic hydrogel. Healthy intervertebral discs, in contrast, retain their ability to hydrodynamically cushion axial compressive forces in the spine over extended periods because living cells within the discs renew the natural hydrogel (i.e., extracellular matrix) component.

Restoration of a clinically useful degree of normal hydrodynamic function in degenerated intervertebral discs is an object of the present invention, and the methods and compositions described herein have been shown to induce and/or enhance such regeneration.

## Disclosure of Invention was a present professor and processor and proces

The present invention comprises methods and compositions for intervertebral disc regeneration. In preferred embodiments, the compositions comprise a three-dimensional fluid matrix of digestion-resistant, cross-linked nucleus pulposus tissue from a donor vertebrate. The donor may be the patient or another animal of the same or different species. Cross-linking of donor nucleus pulposus tissue for the present invention is preferably achieved through use of one or more photooxidative catalysts which selectively absorb visible light. See U.S. patent Nos. 5,147,514, 5,332,475, 5,817,153, and 5,854,397, all incorporated herein by reference. Other

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cross-linking approaches may be used without departing from the scope of the invention, however.

Prior to cross-linking the tissues, chondrocytes of the donor vertebrate are preferably destroyed, fragmented, and/or removed (i.e., decellularized). A preferred decellularization approach involves soaking the tissue in a solution having high concentrations of salt (preferably NaCl) and sugar (preferably sucrose). Such high-salt, high-sugar solutions are referred to as HSHS solutions. Other decellularization approaches may be used, however. After the tissues are decellularized and cross-linked, the resulting fluid matrix may be lyophilized for sterilization and storage, and then rehydrated prior to use. Figure 2 illustrates a process for producing a preferred embodiment of the fluid matrix of the present invention.

The fluid matrix of the present invention is biocompatible, substantially non-immunogenic, and resistant to degradation in vivo. As such, it is capable of providing important internal structural support for an intervertebral disc undergoing regeneration during a period of accelerated proteoglycan synthesis. The cross-linked matrix may be delivered to the intervertebral disc space by injection through a syringe (as depicted in Figure 2), via a catheter, or other methods known in the art.

The three-dimensional fluid matrix of the present invention may be used alone or in combination with growth factors and/or living cells to facilitate regeneration of the structures of a degenerated disc. In patients having sufficient viable endogenous disc cells (chondrocytes) and cell growth factors, the three-dimensional cross-linked matrix alone may substantially contribute to the regeneration of hydrodynamic function in an intervertebral disc in vivo by providing improved mechanical stability of the disc and a more favorable environment for cellular growth and/or metabolism. Conversely, in another embodiment of the invention, a combination of the three-dimensional matrix and one or more purified, preferably bone-derived, cell growth factors may also be used to treat DDD in discs containing viable chondrocytes in a depleted proteoglycan hydrogel matrix. In this case, the cross-linked collagen provides an expanded remodelable threedimensional matrix for the existing (native) chondrocytes within a disc, while the cell growth factors induce accelerated proteoglycan production to restore the hydrogel matrix of the patient. The combination of the three-dimensional matrix and one or more purified cell growth factors is referred to as a cell growth medium. The present invention may also comprise an injectable cell growth medium. Individual purified cell growth factors may be obtained by recombinant techniques known to those skilled in the art, but a preferred plurality of bone-derived purified cell growth factors for the present invention is disclosed in U.S. patent numbers 5,290,763,

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5,371,191 and 5,563,124, all incorporated herein by reference. Bone-derived cell growth factors produced according to these patents are hereinafter referred to as "BP."

Disc regeneration occurs as the cross-linked collagen and proteoglycan matrix supports living cells (which may include exogenous cells as well as native disc or other autologous cells) having inherent capability to synthesize Type II collagen fibrils and proteoglycans in vivo, among other extracellular matrix molecules. Where the patient's native disc cells have been removed or are otherwise insufficient to cause such proliferation, living cells may be added to the three-dimensional matrix of cross-linked nucleus pulposus material to further promote disc regeneration. Accordingly, in another embodiment, the present invention comprises a three-dimensional matrix of cross-linked nucleus pulposus tissue to which exogenous and/or autologous living cells have been added. The injectable combination of three-dimensional matrix material and exogenous and/or autologous living cells is termed herein an injectable cell matrix. Suitable cells for such an injectable cell matrix may be obtained, for example, from the nucleus pulposus of a mammalian vertebral disc, from cartilage, from fatty tissue, from muscle tissue, from bone marrow, or from bone material (i.e., mesenchymal stem cells), but are not limited to these tissue types. These cells are preferably cultured in vitro to confirm their viability and, optionally, to increase the cells' proliferation and synthesis responses using cell growth factors.

Growth factors may optionally be added to cell cultures to stimulate cellular development and elaboration of Type II collagen fibrils and proteoglycans suitable for maintaining an effective disc hydrogel matrix in vivo. An injectable fluid combining purified cell growth factors and a plurality of living cells is termed an injectable cell suspension, and is useful in treating DDD. While an injectable cell matrix alone (i.e., without growth factors) may substantially regenerate hydrodynamic function in an intervertebral disc in vivo if sufficient native cell growth factors are present in the disc, purified (exogenous) cell growth factors may be added to an injectable cell matrix of the present invention to form yet another embodiment of the present invention.

### **Brief Description of Drawings**

Figure 1 is a diagram illustrating components of healthy nucleus pulposus tissue in a vertebrate.

Figure 2 is a diagram illustrating a process for preparation and use of a cross-linked matrix of porcine nucleus pulposus tissue in a preferred embodiment of the invention.

Figure 3 is a photographic reproduction of an SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) analysis comparing the amount of proteins extracted from a cross-linked matrix of the present invention with a non-cross-linked control.

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Figure 4 is a photographic comparison of an H & E (hematoxylin and eosin) stained section of fresh porcine nucleus pulposus tissue with a cross-linked matrix of the present invention, both at 300X magnification.

Figure 5 is a photographic reproduction of a stained nitrocellulose membrane comparing the reactivity of Type II collagen digested from a cross-linked matrix of the present invention and a non cross-linked control.

Figure 6 is a comparison graph of the hydraulic/swelling capacity of a cross-linked matrix of the present invention and a non-crosslinked control.

Figure 7 is a diagram of an experimental process used to demonstrate stimulation of sheep cell ingrowth, proliferation, and new matrix synthesis in an embodiment of the present invention comprising a cross-linked matrix combined with bone protein growth factors (BP).

Figure 8 is a graph and a photograph indicating the results of an Alcian blue assay for matrix production in sheep nucleus pulposus cells stimulated by growth factors.

Figure 9 is a graph indicating the results of immunogenicity tests for a cross-linked matrix of the present invention in rabbit immunizations and sheep serum.

Figure 10 is a diagram of the protocol for an in vivo study of a matrix and growth factor combination of the present invention.

Figure 11 is a radiograph of a vertebral column from a sheep sacrificed at 2 months after an injection of a matrix and growth factor combination in an in vivo study of an embodiment of the present invention.

Figure 12 is a photographic reproduction of histology slides of vertebral discs of a sheep sacrificed at 2 months after an injection of a matrix and growth factor combination of the present invention.

Figure 13 is a radiograph of a vertebral column of a sheep sacrificed at 4 months after an injection of a matrix and growth factor combination in an in vivo study of the present invention.

Figure 14 is a photographic reproduction of histology slides of vertebral discs of a sheep sacrificed at 4 months after an injection of a matrix and growth factor combination of the present invention.

Figure 15 is a graph representing the results of an ELISA performed to measure the synthesis of Type II collagen and Chondroitin-6-sulfate under growth factor stimulation

Figure 16a is a graph indicating the results of an Alcian blue assay for proteoglycan synthesis in human intervertebral disc cells stimulated by growth factor.

Figure 16b is a graph indicating the results of an Alcian blue assay for proteoglycan synthesis in another human intervertebral disc cells stimulated by growth factor.

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Figure 17 is a graph depicting the results of an Alcian blue assay for proteoglycan synthesis in baboon intervertebral disc cells stimulated by growth factor.

Figure 18 is an SDS-PAGE gel of HPLC fractions 27-16 from a sample of BP.

Figure 19 is an SDS-PAGE gel of HPLC fractions 27-16 with identified bands indicated according to the legend of Figure 20.

Figure 20 is an SDS-PAGE gel of BP with identified bands indicated.

Figure 21 is a 2-D (two-dimensional) SDS-PAGE gel with internal standards indicated by arrows.

Figure 22 is a 2-D SDS-PAGE gel with circled proteins (growth factors) identified as in legend.

Figures 23A-23O are Mass Spectrometer results for tryptic fragments.

Figure 24 is a 2-D gel Western blot with anti-phosphotyrosine antibody.

Figures 25A-25D are 2-D gel Western blots with antibodies for the indicated proteins.

For Figure 25A, the growth factors are BMP-3 and BMP-2; for Figure 25B the growth factors are BMP-3 and BMP-7; for Figure 25C the growth factors are BMP-7 and BMP-2; and for Figure 25D the growth factors are BMP-3 and TGF-β1.

Figure 26 is a PAS (periodic acid schiff) stained SDS-PAGE gel of HPLC fractions.

Figure 27 is an anti-BMP-7 stained SDS-PAGE gel of PNGase F treated BP.

Figure 28 is an anti-BMP-2 stained SDS-PAGE gel of PNGase F treated BP.

Figures 29A-29B are bar charts showing explant mass of glycosylated BP samples (Figure 29A) and ALP Score (Figure 29B) of the same samples.

Figure 30 is a chart showing antibody listing and reactivity.

Figures 31A-31B together comprise a chart showing tryptic fragment sequencing data.

Figures 32A-32F together comprise a chart showing tryptic fragment mass spectrometry data.

Figures 33A-33B are an SDS-gel of BP (Figure 33B) and a scanning densitometer scan (Figure 33A).

Figure 34 is a chart illustrating the relative mass of major components of BP.

Best Mode for Carrying Out the Invention

In a preferred embodiment, the invention comprises a biodegradable matrix, which is delivered as an incompressible fluid to induce and/or enhance regeneration or repair of tissues in the intervertebral disc. The biodegradable matrix comprises hydrophilic molecules, which will maintain and/or increase the "captured" water content in intervertebral disc tissues. The

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biodegradable matrix may also serve as a carrier substrate for added growth factors and/or appropriate living cell types.

Since the biodegradable matrix of the present invention is a viscous fluid, it furnishes incompressible support when delivered within a closed, secure disc space. Moreover, because it is distributed uniformly within a disc, the present fluid matrix has a force distribution effect, hydraulically transmitting forces evenly inside the disc. The matrix thus provides resistance against axial compression and annulus collapse, whereas other matrix materials (for example, polymer sponges and collagen sponges) will rapidly collapse under the axial compressive forces within the disc. Solid matrix materials, in contrast, will concentrate forces from end plates directly onto implants, leading to rapid deterioration of implants and/or end plates.

In a preferred embodiment, the biodegradable matrix of the present invention is injectable. Clinical application to a patient can thus be accomplished using minimally invasive techniques, significantly reducing both the cost of treatment and the likelihood of complications relative to procedures such as partial discectomy or vertebral fusion. Similarly, the present invention avoids the requirement for boring a hole into the annulus to implant a prosthetic replacement nucleus pulposus device, such as a relatively solid biodegradable matrix, or to evacuate nucleus tissue to create space for an implanted biodegradable substrate.

The matrix of the present invention is a natural material, preferably prepared from normal, healthy nucleus tissue of animals and/or humans. Accordingly, the matrix is comprised of proteins and matrix molecules especially adapted for efficient hydrodynamic function in intervertebral discs. Such a matrix remains biodegradable under normal circumstances in the presence of specific cellular enzymes, albeit at a slower rate than endogenous disc matrix. It is an important feature of the invention that matrix breakdown products associated with the present invention are digestible by disc cells. In comparison, some matrix materials previously taught (e.g. polyvinyl alcohol) do not break down by physiological processes. In addition, some synthetic polymer substrates create acidic degradation byproducts, in particular PGA and PLA.

Immediate (substantially homogeneous) dispersion of cells within the present matrix is another advantage of the invention. The viscous fluid formulation preferred for injection can be mixed directly with cells of the appropriate type(s) and then delivered immediately to treat an intervertebral disc. In the matrix of the present invention it is not necessary to culture cells and matrix together for some days or weeks before implantation, as it is for certain matrix materials such as PGA and collagen sponges.

The matrix of the present invention is an appropriate substrate for cells, uniquely suited to the ingrowth, proliferation, and residence of intervertebral disc cells. Intervertebral disc cells

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preferentially grow into and survive in the matrix of the present invention, compared to type I collagen sponges fixed with formalin or glutaraldehyde.

The following examples illustrate the preparation of preferred embodiments of the invention and demonstrate its non-immunogenic and disc regenerative properties.

## EXAMPLE 1: Preparation of a Cross-Linked, Fluid Matrix Suitable for Treatment of Degenerative Disc Disease

A three-dimensional fluid matrix of cross-linked nucleus pulposus tissue in accordance with an embodiment of the present invention may be prepared from donor vertebrates. Although porcine donors were used in a particularly preferred embodiment, nucleus pulposus tissues from other vertebrates may also be used, although mammalian vertebrates are preferred (e.g., human, porcine, bovine, ovine, etc.).

Although nucleus pulposus tissues may be harvested by a variety of methods from many vertebral donors, in a preferred embodiment nucleus pulposus tissues were dissected aseptically from spinal intervertebral discs of pigs. In a sterile environment (i.e., a laminar flow hood), the annulus fibrosus of porcine donors was sliced radially and the vertebral end plates separated to expose the nucleus pulposus. The latter material was curetted out of the central portion of the disc, devoid of annulus and end plate tissues.

The nucleus pulposus tissues thus harvested were inserted into sterile dialysis (filter) tubing having a preferred molecular weight cutoff of about 3500 Daltons to substantially prevent loss of low molecular weight proteoglycans from the tissues while substantially reducing bacterial or other contamination. Other semipermeable membranes or filtering membrane types may be used to perform these functions.

The nucleus pulposus tissues to be cross-linked are also preferably treated to destroy and remove donor cells and/or cell fragments. To this end, dialysis tubing containing nucleus pulposus tissues was submerged in a high-salt, high-sucrose (HSHS) solution of about 2.2%: 8.4% w/v (respectively) for about 48 hours. Concentration ranges for the HSHS solution may be from 1% to 50%, but a preferred HSHS solution contains 220 grams NaCl and 837.5 grams of sucrose in 10L water. Preferred HSHS incubation times are from about 24 to about 72 hours, although shorter or longer times may also advantageously be used. Exposure to this HSHS solution results in osmotic destruction and fragmentation of native chondrocyte cells (decellularization), and further results in denaturation of soluble cellular proteins and nucleic acids. The HSHS solution may also contain other reagents which further degrade nucleic acids (including but not limited to sulfones and nucleases), and other reagents which can extract membrane lipids (including but not limited to alcohol, chloroform, and methanol). Although

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native cells of the donor may be retained in other embodiments of the invention, decellularization and denaturation are preferred where exogenous (particularly xenogeneic) tissues are used, so as to reduce the potential for immunogenic responses. Processes other than exposure to HSHS solutions may be used for his purpose.

Cross-linking of the nucleus pulposus tissues is preferably accomplished by a photo-mediated process in accordance with U.S. patent Nos. 5,147,514, 5,332,475, 5,817,153, and/or 5,854,397. In one such process, a photoactive dye (methylene blue) was dissolved in the HSHS solution at a preferred dye concentration of about 20 mg/liter. The photoactive dye was allowed to permeate the nucleus tissues within the dialysis tubing during the initial storage/decellularization process in HSHS. A wide range of photoactive dyes and concentrations, as taught in the foregoing patents, may be used to obtain cross-linked fluid matrices suitable for use in regenerating mammalian disc tissues. Preferred dyes include methylene blue and methylene green at concentrations of about 0.001% to about 1.0% w/v.

To cross-link the collagen within the nucleus tissues, the dialysis tubing containing the dye-permeated nucleus tissues was placed in a photooxidation chamber and exposed to broad-spectrum visible light for 48 hours. In preferred embodiments of the invention, the tissues may be cross-linked from about 24 to about 72 hours. A solution of methylene blue in phosphate buffered saline (PBS) was maintained under controlled temperature at 10°C and circulated around the dialysis tubing within the photooxidation chamber to provide substantially constant temperature regulation of the nucleus tissues. Precise temperature control is not critical to the practice of the invention; however, maintaining a relatively cooler temperature is preferred to avoid damaging the tissues. Following photo-crosslinking of the collagen, the treated nucleus tissues were collected, lyophilized in a vacuum under centrifugation, and finely pulverized in a freezer-mill under liquid nitrogen. The cross-linked matrix product thus prepared can be sterilized using gamma radiation, ethylene oxide (or other sterilants) and stored at -80°C until rehydrated for use. A preferred process for preparing a matrix according to the present invention is illustrated in Figure 2.

In addition to preparation of the cross-linked matrix, control (non-crosslinked) tissues were prepared following the above procedures, except that they were not exposed to light. These control, non-crosslinked tissues were used for comparison purposes.

To investigate the swelling capacity of cross-linked matrix versus non-crosslinked control, lyophilized samples of cross-linked matrix and non-crosslinked control were suspended in water and the increase in weight due to water absorption was measured at various times from 0

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to 96 hours. As illustrated in Figure 6, the cross-linked matrix retained 95% of the hydraulic capacity of the non-crosslinked control.

## EXAMPLE 2: Testing of Fluid Matrix to Evaluate Protein Modification Induced by the Cross-Linking Process

One half gram of the matrix material obtained prior to the lyophilization step of EXAMPLE 1 was placed in 15 mls of a solution of 4M guanidine hydrochloride and agitated on a shaker for 24 hours to solubilize proteoglycans. After centrifugation, the supernatant was discarded and the pellet washed in distilled water 3 times for 5 minutes each. The pelleted matrix material was then removed and blot-dried on filter paper.

One hundred mg of the blot-dried matrix was placed in a 1.5 ml microcentrifuge tube with 1000 µl of 1% sodium dodecyl sulfate (SDS) containing 5% beta-mercaptoethanol (BME). The matrix in SDS/BME was boiled for one hour to extract proteins (e.g., collagens). Samples were then centrifuged at 12000 rpm for 1 hour and aliquots of the supernatant were subjected to electrophoresis in gradient polyacrylamide gels.

Gels were stained with Coomassie blue or silver to visualize proteins extracted by the SDS/BME and heat treatment. As illustrated in Figure 3, collagen bands stained prominently in control, non-crosslinked tissues but exhibited only faint staining in cross-linked matrix. These results demonstrated that in the cross-linked matrix material, collagen proteins were not easily extracted by the above treatment, indicating that crosslinking had occurred. In contrast, stained gels of the control tissues demonstrated that collagen proteins were readily extracted from non-crosslinked material by the above treatment. See Figure 3.

## EXAMPLE 3: Matrix Histology to Evaluate Cellular Debris and Residual Membranous Material

Cross-linked matrix material obtained prior to the lyophilization step of Example 1 was placed in 4% paraformaldehyde for tissue fixation. Standard histology techniques of embedding, sectioning, and staining of sections with hematoxylin & eosin dyes were performed. Visualization of cross-linked matrix in H & E-stained sections demonstrated that the matrix preparation process facilitates destruction of cellular membranes and intracellular elements, with minimal membrane material remaining as compared to fresh porcine nucleus pulposus material as well as non-crosslinked tissue decellularized by HSHS treatment, freeze-thaw cycles, and HSHS treatment plus freeze-thaw cycles. See Figure 4

# EXAMPLE 4: Evaluation of Matrix Antigenic Reactivity Using Monoclonal Antibodies to Type II Collagen

Cross-linked matrix material obtained prior to the lyophilization step of Example 1 was also subjected to pepsin digestion to cleave Type II collagen proteins. The protein digests were

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run on SDS/PAGE and then transferred to a nitrocellulose membrane. Total protein transferred to the membrane was visualized using Colloidal Gold.

The visualized nitrocellulose membranes were incubated with a mouse monoclonal antibody to Type II collagen and a secondary antibody (anti-mouse) conjugated with alkaline phosphatase. The antibody reactivity was visualized through addition of alkaline phosphatase substrate. As depicted in Figure 5, the antibodies toward Type II collagen did not react with pepsin digests of the cross-linked matrix as much as with the pepsin digests of the non-crosslinked control tissue. The results indicate that the matrix of the invention may have reduced antigenic epitopes for Type II collagen, and thus have less immunogenicity than non-crosslinked tissues. See Figure 5.

#### EXAMPLE 5: Evaluation of Matrix Immunogenicity in Rabbit Antisera Production

One gram of the lyophilized and pulverized matrix material prepared according to EXAMPLE 1 was dispersed in PBS (i.e., rehydrated) and centrifuged. The protein concentration of the supernatant was then determined using the BCA assay and the supernatant was diluted with PBS to a final concentration of 200 µg of protein per ml of PBS. The diluted supernatant was then sterilized for injection protocols. Three rabbits were immunized with 100 µg of protein from the sterilized supernatant. Each rabbit received 9 immunizations over a 14 week period and sera was collected from the rabbits on a regular schedule.

Antisera production against the protein extract was measured using an enzyme-linked immunosorbent assay (ELISA). Type II collagen was included as a positive control in the ELISA. Colorimetric evaluation of antisera directed against the matrix material demonstrated very low immunogenicity in rabbits. See Figure 9.

## EXAMPLE 6: Matrix Formulation Including Serum and Other Fluids For Injections And Delivery

One gram of the lyophilized and pulverized matrix material prepared according to EXAMPLE 1 was sterilized with 70% ethanol and the ethanol was removed by successive PBS rinses. The dispersed matrix was centrifuged and the pellet was suspended in heat-inactivated sheep serum at a ratio of 0.5g lyophilized matrix to 1 ml serum to prepare a viscous fluid matrix which can be loaded into a standard syringe and delivered via a small gauge needle. In preferred embodiments of the invention, the serum is collected from the vertebrate animal or human patient to be treated, heat-inactivated to destroy unwanted protein components (complement proteins), and passed through a 0.2 micron sterile filtration unit. Different matrix/serum ratios may also be advantageously employed. Ratios ranging from 0.1g to 2.0 g of lyophilized matrix to 1 ml of The control in serum are preferred. The control is a series of the control in the

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Serum is a preferred fluid for mixture and delivery of the cross-linked matrix of the present invention because it contains various intrinsic growth factors that are beneficial to intervertebral disc cells. Serum also serves as a suitable carrier for extrinsic protein growth factors and/or small molecules. The beneficial effects of extrinsic growth factors on intervertebral disc cells are enhanced by the addition of serum.

Other fluids are also suitable for mixture and delivery of the viscous fluid matrix. For example, sterile saline or sterile water may also be used. The examples herein are not meant to be limiting as to the variety of carrier fluids which may be used to mix and deliver the matrix in the present invention.

# EXAMPLE 7: Injection of Matrix Formulation To Intervertebral Discs Using Pressure-Mediated Syringe

Matrix material was prepared according to EXAMPLE 6 (mixed with serum) to form a viscous fluid and loaded into a standard syringe having a small gauge needle (e.g., 18-31 gauge) attached. Syringe injection pressure can be controlled simply by the fingers of the hand. In other embodiments of the invention, pressure to inject the viscous fluid can be controlled by an external device which concomitantly measures (e.g., via a pressure transducer) and delivers (e.g., by compressed air) a predetermined force to the syringe plunger.

In one preferred embodiment of this device, a thermal element is included in the needle. By providing a needle having a thermal element, it is possible to deliver heat to the outer layers of the annulus fibrosus at the end of the treatment and during removal of the syringe needle in order to shrink collagen fibers around the needle and effectively seal the site of needle penetration.

It is further contemplated that the matrix of the present invention can be delivered to the disc space of a patient transpedicularly (i.e., through the pedicle of the vertebrae). In particular, the cross-linked matrix can be administered percutaneously via a biopsy cannula inserted through a channel in the pedicle. After delivery of the matrix, the channel can then be filled with bone cement or other like material to seal the channel.

EXAMPLE 8: Isolation of Human, Sheep, and Baboon Intervertebral Disc Nucleus Pulposus
Cells

Human intervertebral nucleus pulposus tissues were collected during surgery, suspended in Dulbecco's Modified Eagle Medium/ Nutrient Mixture F-12 (DMEM/F-12) in a 1:1 v/v mixture supplemented with antibiotics. The tissues were kept on ice until dissection, at which time they were rinsed 2-3 times in sterile Dulbecco's Phosphate Buffer Saline (DPBS) to remove any blood. In a laminar flow hood, the nucleus tissues were isolated and diced into small (2 mm)

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cubes, and then placed in a Tissue Culture Medium (hereinafter referred to as "TCM") comprising DMEM/F-12 culture media supplemented with 10% heat inactivated fetal bovine serum, 0.25% penicillin, 0.4% streptomycin, 0.001% amphotericin B, and 50µg/ml ascorbic acid. Only tissues clear of blood and other anomalous elements were used. Placed on a shaker at 37oC, the tissues were digested with 0.01% hyaluronidase (Calbiochem) in TCM for 2 hours, 0.01% protease (Sigma) in TCM for 1 hour, and 0.1% collagenase Type II (Sigma) in TCM overnight to obtain a suspension of human intervertebral disc nucleus pulposus cells.

The foregoing procedure was also applied to sheep and baboon intervertebral disc nucleus pulposus tissues to obtain suspensions of sheep and baboon intervertebral disc nucleus pulposus cells, respectively.

EXAMPLE 9: Primary Culture and Expansion of Human, Sheep, and Baboon Intervertebral Disc Nucleus Pulposus Cells

Human intervertebral disc nucleus pulposus cells from EXAMPLE 8 were expanded by culturing in TCM at 37oC in 5% CO2 atmosphere and 95% relative humidity. The TCM was changed every 2-3 days and the cells were passaged with trypsin to another container, when 80-90% confluent, for continued expansion.

The foregoing procedure was also applied to sheep and baboon intervertebral disc nucleus pulposus tissues to obtain an expanded supply of sheep and baboon intervertebral disc nucleus pulposus cells.

20 EXAMPLE 10: Alcian Blue Assay of Disc Cell Matrix Production in Human, Sheep, and Baboon Intervertebral Disc Nucleus Pulposus Cells

Human intervertebral disc cells from EXAMPLE 9 were seeded and grown in 24 well plates in TCM in the presence or absence of exogenous growth factors. At various time points, TCM was aspirated out from the wells and the wells washed 3 times with PBS. The cells were then fixed with 4% paraformaldehyde (pH 7.4) for 10min. The fixed cells were washed 2 times with PBS and then stained overnight with 0.5% Alcian blue in 0.1N hydrochloric acid (pH 1.5). After overnight staining, excess stain was rinsed out with 3 rinses of PBS. The remaining Alcian blue stain (bound to proteoglycans) was dissolved overnight into 6M guanidine hydrochloride and the absorbance at 630nm was measured using a spectrophotometer, providing an indication of the induction of matrix production by exogenous growth factors in human nucleus pulposus cells.

The foregoing procedure was also applied to sheep and baboon intervertebral nucleus pulposus cells from EXAMPLE 9 to obtain an indication of the induction of matrix production by exogenous growth factors in sheep and baboon nucleus pulposus cells.

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## EXAMPLE 11: Enzyme Linked Immunosorbent Assay (ELISA) on Ovine Intervertebral Disc Nucleus Pulposus Cells

To detect specific antigenic epitopes in the synthesized matrix, sheep intervertebral nucleus pulposus cells from EXAMPLE 9, seeded and grown in monolayer, were fixed in 2% glutaraldehyde for 1 hour at room temperature. The fixed cells were washed 3 times with TBS for 5 min. each. To block non-specific antibody binding, the cells were incubated in a solution of Tris buffered saline (TBS) containing 1mM ethylene-diamine-tetraacetic acid (EDTA), 0.05% Tween-20, and 0.25% bovine serum albumin for 1hour. The blocking step was followed by 3 washes with TBS for 5 min. each. The cells were incubated with the primary antibody at room temperature for 2.5 hours, and the excess primary antibody was removed by 3 washes with TBS for 5 min. each. A second incubation with blocking buffer was performed for 10 min., followed by 3 washes with TBS. The cells were then incubated with the secondary antibody, which was conjugated with alkaline phosphatase enzyme, for 3 hours at room temperature. The unbound secondary antibodies were removed by 3 washes of TBS for 5 min each. The bound primary and secondary antibodies were detected by addition of an enzyme-specific substrate which produced a colored reaction. The colorimetric measurement was performed using a spectrophotometer, providing a quantitative measure of the presence of the bound antibodies.

# EXAMPLE 12: Effect of Exogenous Growth Factors on Proteoglycan Synthesis in Ovine Intervertebral Disc Nucleus Pulposus Cells

Transforming growth factor-μ1 (TGFβ1) and a mixture of bone-derived protein growth factors (BP) produced according to U.S. patent Nos. 5,290,763, 5,371,191 and 5,563,124, were tested for their effects on stimulation of proteoglycan synthesis in ovine nucleus pulposus cells. Sheep intervertebral disc nucleus cells were collected and cultured as described in EXAMPLES 8 and 9. Sheep cells were seeded in micromass (200,000) into the wells of a 24 well plate. Growth factor dilutions were prepared in TCM supplemented with 0.5% heat-inactivated fetal bovine serum. TGFβ1 and BP were both tested at 10 ng/ml; BP was also tested at a concentration of 10 μg/ml. Control wells without growth factors contained TCM supplemented with 0.5% and 10% heat-inactivated fetal bovine serum. The cells were incubated in continuous exposure to the various growth factors for 7 and 10 days. At these time points, the cells were fixed and the amount of proteoglycan synthesis was measured by the Alcian blue assay as described in EXAMPLE 10.

At both 7 and 10 day time points, proteoglycan synthesis was significantly greater in the 10% fetal bovine serum control cultures than in the 0.5% fetal bovine serum control cultures. At the 7 day time point, BP at the higher 10 µg/ml concentration produced a significant (93%)

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increase in proteoglycan synthesis above the level in 10% serum control culture and a greater (197%) increase above the 0.5% serum control. Slight increases in proteoglycan synthesis above the 0.5% serum control were observed in the 10 ng/ml TGF $\beta$ 1 and BP cultures, but these increases were not significant.

At the 10 day time point (Figure 8), 10  $\mu$ g/ml BP produced a significant increase (132%) in proteoglycan synthesis over the 10% serum control, while 10 ng/ml TGF $\beta$ 1 produced a significant increase (52%) above the 0.5% serum control. At 10 ng/ml, BP exhibited a modest 20% increase in proteoglycan synthesis over the 0.5% serum control, while at the 10  $\mu$ g/ml concentration, BP produced an 890% increase above the 0.5% serum control.

# EXAMPLE 13: Effect of Exogenous Growth Factors on Type II Collagen and Chondroitin-6-Sulfate Produced by Ovine Intervertebral Disc Nucleus Pulposus Cells

TGF $\beta1$  and BP were tested for their effects on stimulation of Type II collagen and chondroitin-6-sulfate synthesis in sheep intervertebral disc nucleus pulposus cells. The cells were obtained and cultured as described in EXAMPLES 8 and 9 and seeded into tissue culture dishes. The TGF $\beta1$  and BP growth factors were prepared in TCM supplemented with 0.5% heat inactivated fetal bovine serum. TGF $\beta1$  was tested at a concentration of 10 ng/ml; BP was tested at a concentration of 10  $\beta$ g/ml. Control cultures were incubated in TCM supplemented with 0.5% serum alone.

After incubation with growth factors for 7 days, cell cultures were fixed in 2% glutaraldehyde and the quantity of Type II collagen and chondroitin-6-sulfate produced in the cell cultures was detected by ELISA according to the procedure described in EXAMPLE 11. The primary antibodies used were mouse anti-human Type II collagen and mouse anti-human chondroitin-6-sulfate.

At 7 days, cell cultures incubated with 10  $\mu$ g/ml BP produced 324% more Type II collagen and 1780% more chondroitin-6-sulfate than control cultures. 10 ng/ml TGF $\beta$ 1 increased production of Type II collagen by 115% and chondroitin-6-sulfate by 800% over controls. See Figure 15.

# EXAMPLE 14: Effect of Exogenous Growth Factors on Proteoglycan Synthesis in Human Intervertebral Disc Nucleus Pulposus Cells

TGFβ1 and BP were tested for their effects on stimulation of proteoglycan synthesis in human nucleus cells. Human intervertebral disc nucleus pulposus cells obtained from Disc L5-S1 of a 40 yr old female patient were cultured as described in EXAMPLES 8 and 9 and seeded into 24 well plates. After the cells adhered to the well surface, multiple dilutions of different growth factors were added. The concentrations of growth factors tested were 10 ng/ml TGFβ1, and 10

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and 20 µg/ml of BP. The dilutions were prepared in TCM. The cells were fixed after 5 and 8 days of continuous exposure to growth factors and proteoglycans synthesized were detected by the Alcian blue assay as described in EXAMPLE 10.

At 5 days only BP produced a significant increase in Alcian blue staining over controls. At 10  $\mu$ g/ml BP there was a 34% increase over the control while at 20 $\mu$ g/ml there was a 23% increase over the control. The difference between the averages of 10 and 20  $\mu$ g/ml BP was not significant.

At 8 days (Figure 16a), both growth factors exhibited a significant increase in Alcian blue staining over the control. TGF $\beta$ 1 at 10ng/ml had a 42% increase over the control. BP had a 60% increase at 10 $\mu$ g/ml and 66% increase at 20 $\mu$ g/ml over the control.

# EXAMPLE 15: Effect of Exogenous Growth Factors on Proteoglycan Synthesis in Human Intervertebral Disc Nucleus Pulposus Cells

A second experiment to test the effects of TGFβ1 and BP on proteoglycan synthesis was performed on a different human patient from that described in EXAMPLE 14. Human intervertebral disc cells obtained from another 40-year-old female patient were cultured as described in EXAMPLES 8 and 9 and seeded into 24 well plates. Growth factors were added after the cells were allowed to adhere overnight. TGFβ1 was tested at a concentration of 10 ng/ml; BP was tested at 10 μg/ml. After 6 and 9 days the cells were fixed and the amount of proteoglycans synthesized was measured by the Alcian blue assay as described in EXAMPLE 10.

At 6 days cells stimulated with 10 ng/ml TGF $\beta$ 1 produced 54% more proteoglycans than control, and 10  $\mu$ g/ml BP increased production by 104% over the control. At 9 days (Figure 16b), 10 ng/ml TGF $\beta$ 1 increased production by 74% over controls, and 10  $\mu$ g/ml BP increased production by 171% over the control.

# EXAMPLE 16: Effect of Exogenous Growth Factors on Proteoglycan Synthesis in Baboon Intervertebral Disc Nucleus Pulposus Cells

TGFβ1 and BP were tested for their effects on stimulation of proteoglycan synthesis in baboon nucleus cells. Baboon intervertebral disc nucleus pulposus cells were obtained from a 7 year old male baboon, cultured as described in EXAMPLES 8 and 9, and seeded into a 24 well plate. The cells were allowed to adhere to the well surface before the addition of growth factors.

The concentrations of growth factors tested were 10 µg/ml BP and 10 ng/ml TGFβ1. The dilutions were prepared in TCM. The cells were fixed after 4 and 8 days of continuous exposure to growth factors, and proteoglycan synthesis was detected by the Alcian blue assay as described in EXAMPLE 10.

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At 4 days there was no significant increase in proteoglycan synthesis between the different growth factors and the control. At 8 days (Figure 17), TGFβ1 and BP significantly increased proteoglycan synthesis over the control, but the increase was only marginal. In particular, TGFβ1 produced a 21% increase over the control while BP produced a 22% increase over the control.

## EXAMPLE 17: Staining of Seeded Matrix Material with Phalloidin

Cross-linked matrix seeded with living cells was stained with phalloidin to indicate the growth and proliferation of living cells into the matrix. The media was rinsed from the matrix with 3 PBS washes of 5 min each. The matrix was fixed for 1 hour at room temperature with 4% paraformaldehyde. The 4% paraformaldehyde was washed off with 3 PBS rinses. The matrix was treated with 0.1% Triton-X 100 for 3min and then washed with 3 PBS rinses. The matrix was then stained with phalloidin-conjugated rhodamine, made up in PBS, for 45 min. Excess phalloidin was washed off with PBS. The matrix was mounted on slides and viewed under fluorescence with filter of  $\lambda$  range 530-550 nm.

# 15 EXAMPLE 18: Growth and Proliferation of Sheep Intervertebral Disc Nucleus Pulposus Cells into Non-Homogenized Matrix with BP Growth Factor

Ingrowth and proliferation of growth factor stimulated sheep intervertebral disc nucleus pulposus cells into the matrix of the present invention was investigated. Cross-linked matrix material obtained prior to the lyophilization step of Example 1 was cut into square pieces 75mm on each side and sterilized in 70% ethanol for 3 hours. Remaining steps in the protocol were performed under aseptic conditions.

Ethanol was removed from the matrix with two 1-hour washes in sterile PBS, followed by a one hour wash in TCM. The matrix pieces were then suspended overnight in TCM having BP concentrations of 20ng/ml and 20μg/ml. The control was cross-linked matrix suspended in 20 μg/ml BSA (bovine serum albumin). Each matrix piece was then placed in a well of a 24 well plate and seeded with TCM containing sheep intervertebral disc nucleus cells at 40,000 cells/ml. The cells were allowed to grow into the matrix and the TCM was changed every 2-3 days. Sample matrix pieces were fixed at 3, 6 and 9 days and stained with phalloidin as described in EXAMPLE 17. The process is illustrated in Figure 7.

Infiltration of sheep nucleus pulposus cells into the matrix was observed at all of the 3, 6 and 9 day timepoints, indicating that the matrix is biocompatible. The number of cells observed per field was higher at 6 and 9 days, indicating that the cells were proliferating into the matrix. More cells were observed in matrix pieces that had been suspended in TCM containing BP than

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in controls having no growth factor. BP at 20  $\mu$ g/ml produced the greatest infiltration and proliferation of cells into the matrix.

EXAMPLE 19: Growth and Proliferation of Sheep Intervertebral Disc Nucleus Pulposus Cells into Homogenized Matrix with BP Growth Factor

A further investigation of the ingrowth and proliferation of growth factor stimulated sheep intervertebral disc nucleus pulposus cells into the matrix of the present invention was made using homogenized matrix, as opposed to the non-homogenized matrix in EXAMPLE 18. Cross-linked matrix material obtained prior to the lyophilization step of Example 1 was homogenized using a tissue homogenizer, and sterilized in 70% ethanol for 3 hours. All subsequent steps in the protocol were under aseptic conditions.

The homogenized matrix was centrifuged at 3200 rpm for 10 min and the supernatant was discarded. The pelleted matrix was rinsed with two 1-hour PBS washes, followed by a 1-hour TCM wash. Between each wash the matrix was centrifuged, and the supernatant was discarded. The pelleted matrix was then suspended overnight in TCM having BP concentrations of 20 ng/ml and 20  $\mu$ g/ml. The control was cross-linked matrix suspended in 20  $\mu$ g/ml BSA

The TCM/matrix mixture was then centrifuged and the supernatant was discarded. The matrix pellet was resuspended in TCM containing sheep intervertebral disc nucleus cells, obtained according to the procedure in EXAMPLES 8 and 9. The matrix/cell suspension was pipetted into wells of a 24 well plate. The TCM was changed every 2-3 days. The homogenized matrix seeded with cells was fixed at 4 days and stained with phalloidin as described in EXAMPLE 17. The process is illustrated in Figure 7.

After 4 days, the layer of cross-linked matrix soaked in 20 µg/ml BP and seeded with cells had contracted to form a rounded clump of compact tissue. This tissue was comprised of both the original cross-linked matrix and the newly synthesized matrix produced by the infiltrated cells. There were very few cells adherent to the well surface, indicating that most cells had infiltrated the matrix. This conclusion was reinforced by the dense infiltration of cells into the matrix as visualized by phalloidin staining. The cells had assumed a rounded morphology which is characteristic of nucleus chondrocytic cells, indicating reversion to their original morphology. Cells had also grown into matrix soaked in 20 ng/ml BP by 4 days, but cell ingrowth was not as dense as in the matrix soaked in 20 µg/ml BP.

The control matrix suspended in BSA also had cells infiltrating into it, but it was the least populated among the different dilutions.

# EXAMPLE 20: In Vivo Evaluation of Cross-linked Matrix and Bone Protein (BP) Growth Factor for Nucleus Pulposus Regeneration in an Ovine Lumbar Spine Model

Pilot studies were conducted to evaluate preparative and surgical methods for the implantation of the cross-linked matrix containing BP growth factors into the intervertebral disc space of the sheep lumbar spine, to evaluate whether implantation of the matrix with growth factors arrests degeneration and/or stimulates regeneration of nucleus pulposus in a sheep disc degeneration model over a period of six months, and to assess the antibody- and cell-mediated immune response in sheep to the matrix/BP combination.

#### Study #1

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One-half gram (0.5 g) of cross-linked, lyophilized and pulverized matrix prepared as described in EXAMPLE 1 was rehydrated and sterilized by two 4 hour rinses in 70% isopropanol. The matrix was centrifuged and pelleted, and then rinsed in sterile PBS three times for 2 hours each to remove the isopropanol. The rehydrated matrix was again centrifuged and pelleted.

Bone Protein (BP) prepared according to U.S. patent Nos. 5,290,763 and 5,371,191 was obtained from Sulzer Biologics, Inc. (Wheat Ridge, CO) in a lyophilized form. Two milligrams (2 mg) of BP was suspended in 100 (I dilute 0.01M hydrochloric acid to produce a 20 mg/ml BP stock solution. The BP stock solution was diluted to 100 µg/ml in sheep serum and the BP/serum suspension was sterile-filtered through a 0.2 micron filter. Next, 1.0 ml of the sterile BP/serum suspension was added to 1.0 ml of the rehydrated matrix described above to obtain a final concentration of 50 µg BP per ml of cross-linked, rehydrated matrix/serum suspension. At the time of surgery, one aliquot (0.5 ml) of the rehydrated matrix/BP/serum suspension was loaded into a sterile 3 ml pressure control syringe with an 18 or 20 gauge needle for injection.

Three sheep were anesthetized and the dorsolateral lumbar area prepared for surgery. Blood was drawn from each sheep pre-operatively, centrifuged, and serum collected for immunology studies. A ventrolateral, retroperitoneal approach was made through the oblique abdominal muscles to the plane ventral to the transverse processes of the lumbar spine. The annuli fibrosi of intervertebral discs L3-4, L4-5, and L5-6 were located, soft tissues retracted, and a discrete 5 mm deep by 5 mm long incision was made into both L3-4 and L5-6 discs. The intervening, middle L4-5 disc remained intact to serve as an intra-operative control. Following annulus stab procedures, the musculature and subcutaneous tissues were closed with absorbable suture. After postoperative recovery, sheep were allowed free range in the pasture.

Two months after the annulus stab surgical procedures, the sheep were operated upon a second time. After anesthesia and preparation for surgery, the three operated lumbar spine levels

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were again exposed. Two hundred microliters (200 µl) of the prepared test material (i.e., rehydrated matrix/BP/serum suspension) was injected into the intradiscal space of one (L5-6) of the experimentally-damaged discs. The second operated disc (L3-4) served as a sham-treated degenerative disc; the syringe needle punctured the annulus but no material was injected. After disc treatments, the musculature and subcutaneous tissues were closed with absorbable suture. Following postoperative recovery, sheep were allowed free range of movement. The study design is diagrammatically represented in Figure 10.

The sheep were sacrificed at 2, 4, and 6 months after the second surgery. The radiograph from the 2 month sheep showed a degenerative appearance of the untreated disc but a normal appearance in the control and treated discs (Figure 11). Histological analysis of the 2 month sheep as illustrated in Figure 12 confirmed extensive degeneration within the sham-treated, stab-induced degenerative disc. In both the control disc and the matrix/BP-treated disc, a normal sized gelatinous nucleus and regular, compact annulus were observed. In the 4 month and 6 month sheep, no obvious changes were seen in the radiograph of the three discs. A radiograph of the 4 month sheep is shown in Figure 13. However, on gross dissection in the 4 month sheep, the sham-treated disc exhibited obvious gross degeneration while the control and treated discs were normal in appearance (Figure 14). In the 6 month sheep, there were no gross differences between the sham-treated, control, and treated discs.

Although there was some variation in the rate of degeneration using the annulus stab technique (i.e., the absence of clear degeneration in the 6 month sheep), these results suggest that the cross-linked matrix/BP treatment may protect against or impede the progress of stab-induced degeneration in sheep intervertebral discs.

#### Study #2

For the second study, matrix material was rehydrated and combined with BP and serum to produce a matrix/BP/serum suspension as described in Study #1.

Twelve sheep were anesthetized and the dorsolateral lumbar area prepared for surgery. Blood was drawn from each sheep pre-operatively, centrifuged, and serum collected for immunology studies. A ventrolateral, retroperitoneal approach was made through the oblique abdominal muscles to the plane ventral to the transverse processes of the lumbar spine. The annuli fibrosi of intervertebral discs L1-2, L2-3, L3-4, L4-5, and L5-6 were located, soft tissues retracted, and a small diameter hole punched through the annulus using a syringe needle in 4 of the 5 discs. A small curette was then placed through the hole into the intradiscal space to remove a discrete portion of nucleus pulposus from each of the four discs in each sheep. In 2 of the 4 damaged discs, 0.5 ml of the matrix/BP/serum suspension was injected into the intradiscal spaces

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and the needle punctures were sealed off with ligament sutured over them. The immediate injection of this suspension was considered an "acute" treatment protocol. The 2 other damaged discs were left untreated at that time but were sealed off with ligament sutured over the needle punctures. The intervening, middle L3-4 disc remained intact in all sheep spines to serve as an intra-operative control. Following these procedures, the musculature and subcutaneous tissues were closed with absorbable suture. After postoperative recovery, sheep were allowed free range.

Six weeks after the first surgery to remove portions of the nucleus pulposus, the sheep were operated upon a second time. After anesthesia and preparation for surgery, the five operated lumbar spine levels were again exposed. In one of the two remaining nontreated discs which had been damaged six weeks before, 0.5 milliliters of the prepared test material (i.e., rehydrated matrix/BP/serum suspension) was injected into the intradiscal space of the disc. The injection of this suspension six weeks later into a damaged disc was considered a "delayed" treatment protocol. The second nontreated damaged disc served as a sham-treated degenerative disc; the syringe needle punctured the annulus but no material was injected. The treatment method used in each of the four experimentally-damaged discs was randomized for location within the spines. That is, except for the intact control disc (L3-4), the locations of an "acute" treatment disc, a "delayed" treatment disc, or a nontreated, damaged disc, were randomly assigned to one of the four different lumbar disc levels. After disc treatments, the musculature and subcutaneous tissues were closed with absorbable suture. Following postoperative recovery, sheep were allowed free range.

The sheep were sacrificed at 2, 4, and 6 months after matrix/BP/serum injections and the spines were fixed for histology in formalin. Cross-sections were taken from plastic-embedded discs, stained with H & E and Saffranin-O, and evaluated for chondrocyte proliferation (cloning), proteoglycan staining intensity, level of fibrosis, and level of ossification. An evaluation of the "acute" treatment discs, "delayed" treatment discs, sham-treated, and control discs was made in a blinded fashion and ranked +1, +2, or +3 (low, medium, or high) for each parameter listed above. Semiquantitative evaluation of the histological results was compared in 2 month, 4 month, and 6 month sheep for both the "acute" and "delayed" (6 week) treatments.

The results demonstrated overall that injected matrix # BP stimulated chondrocyte cloning and accumulation of Saffranin-O staining of glycosaminoglycans in the nucleus matrix of damaged discs. In particular, the extent of regenerative repair was much greater in both "acute" treatment discs and "delayed" treatment discs, compared to that observed in non-treated, damaged discs. This greater level of repair in matrix/BP-treated discs was statistically significant at the

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0.01 level of confidence. There was also less fibrosis and ossification seen in the acute and delayed treatment discs compared to the non-treated discs.

A significant difference was also noted between the "delayed" treatment discs and the "acute" treatment discs in the level of proteoglycan staining. For example, Saffranin-O staining as an index to proteoglycan synthesis and content in the nucleus matrix was greater in the "delayed" matrix/BP-treatment discs than in the "acute" matrix/BP-treatment discs. Additional benefits apparent in the histological evaluation, which were associated with "delayed" treatment with matrix/BP, were an overall lack of bony transformation (ossification) or fibrous tissue accumulation (fibrosis) within the treated discs compared to the non-treated, damaged discs. In general, the results in Study #2 support and elaborate earlier indications from Study #1 that treatment of damaged discs with the cross-linked matrix/BP may protect against or impede the progress of degeneration in experimentally-damaged sheep intervertebral discs.

#### EXAMPLE 21: Characterization of BP

Specific growth factors present in the mixture of growth factors produced according to U.S. patent Nos. 5,290,763, 5,371,191, and 5,563,124 (i.e., BP) have been identified. BP has been partially characterized as follows: HPLC fractions have been denatured, reduced with DTT (dithiothreitol), and separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). One minute high performance liquid chromatography (HPLC) fractions taken at from 27 to 36 minutes are shown in Figure 18. Size standards (ST) of 14, 21, 31, 45, 68 and 97 kDa were obtained as Low Range size standards from BIORAD(tm) and are shown at either end of the Coomassie blue stained gel (Figures 18 and 19). In the usual protocol, HPLC fractions 29 through 34 are pooled to produce BP (see box in Figures 18 and 19), as shown in a similarly prepared SDS-PAGE gel in Figure 33B.

An SDS-PAGE gel of BP was also analyzed by Western immunoblot with a series of antibodies, as listed in Figure 30. Visualization of antibody reactivity was by horse radish peroxidase conjugated to a second antibody and using a chemiluminescent substrate. The reactivities are as indicated in Figure 30.

The BP was further characterized by 2-D (two dimensional) gel electrophoresis, as shown in Figures 21 and 22. The proteins are separated in horizontal direction according to charge (pI) and in the vertical direction by size according to the method of O Farrell et al. (Cell, 12:1133-1142; 1977). Internal standards, specifically tropomyosin (33 kDa; pI 5.2) and lysozyme (14.4 kDa, pI 10.5-11.0), are included and the 2-D gel was visualized by Coomassie blue staining. Figure 21 shows the stained 2-D gel with size standards indicated on the left. Tropomyosin (left arrow) and lysozyme (right arrow) are also indicated.

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The same gel is shown in Figure 22 with several identified proteins indicated by numbered circles. The proteins were identified by mass spectrometry and amino acid sequencing of tryptic peptides, as described below. The identity of each of the labeled circles is provided in the legend of Figure 22.

The various components of the BP were characterized by mass spectrometry and amino acid sequencing of tryptic fragments where there were sufficient levels of protein for analysis. The major bands in the 1-D (one dimensional) gels were excised, eluted, subjected to tryptic digestion, purified by HPLC and sequenced by methods known in the art. The major bands are identified by band number, as shown in Figures 19 and 20. The sequence data was compared against known sequences, and the fragments are identified as shown in Figure 31. In some cases, the identification is tentative due to possible variation between the human and bovine sequences and/or possible post translational modifications, as discussed below.

The same tryptic protein fragments were analyzed by mass spectrometry and the mass spectrograms are shown in Figures 23A-23O. The tabulated results are shown in the Table depicted in Figures 32A-32F, which provides identification information for each of the indicated bands, as identified in Figures 19 and 20. As above, assignment of band identity may be tentative based on species differences and post translational modifications.

The identified components of BP were quantified as shown in Figures 33A and 33B. Figure 33B is a stained SDS-PAGE gel of BP and Figure 33A represents a scanning densitometer trace of the same gel. The identified proteins were labeled and quantified by measuring the area under the curve. These results are presented in Figure 34 as a percentage of the total peak area.

As Figure 34 indicates, there are 11 major bands in the BP SDS-PAGE gel representing about 60% of the protein in BP. Further, TGF-β1 was quantified using commercially pure TGF-β1 as a standard, and was determined to represent less than 1% of the BP protein. The identified proteins fall roughly into three categories: the ribosomal proteins, the histones, and growth factors, including active growth factors comprising members of the TGF-β superfamily of growth factors, which includes the bone morphogenic proteins (BMPs). It is believed that the ribosomal proteins and histone proteins may be removed from the BP without loss of activity, and the specific activity is expected to increase correspondingly.

Because several of the proteins migrated at more than one size (e.g., BMP-3 migrating as 5 bands) investigations were undertaken to investigate the extent of post-translational modification of the BP components. Phosphorylation was measured by anti-phosphotyrosine immunoblot and by phosphatase studies. Figure 24 shows a 2-D gel, electroblotted onto filter paper and probed

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with a phosphotyrosine mouse monoclonal antibody by SIGMA (# A-5964). Several proteins were thus shown to be phosphorylated at one or more tyrosine residues.

Similar 2-D electroblots were probed with BP component specific antibodies, as shown in Figures 25A-D. The filters were probed with BMP2, BMP-3 (Fig. 25A), BMP-3, BMP-7 (Fig. 25B), BMP-7, BMP-2 (Fig. 25C), and BMP-3 and TGF-β1 (Fig. 25D). Each shows the characteristic, single-size band migrating at varying pI, as is typical of a protein existing in avarious phosphorylation states.

Native and phosphatase treated BP samples were also assayed for morphogenic activity by explant mass and ALP (alkaline phosphatase) score. The results showed that AcP treatment reduces the explant mass and ALP score from 100% to about 60%.

The BP was also analyzed for glycosylation. Figure 26 shows an SDS-PAGE gel stained with periodic acid schiff (PAS) - a non-specific carbohydrate stain, indicating that several of the BP components are glycosylated (starred protein identified as BMP-3). Figures 27 and 28 show two specific proteins (BMP-7, Figure 27 and BMP-2, Figure 28) treated with increasing levels of PNGase F (Peptide-N-Glycosidase F), and immunostained with the appropriate antibody. Both BMP-2 and BMP-7 show some degree of glycoslyation, but appear to have some level of protein that is resistant to PNGase F, as well (plus signs indicate increasing levels of enzyme). Functional activity of PNGase F and sialadase treated samples were assayed by explant mass and by ALP score, as shown in Figure 29A and 29B, indicating that glycosylation is required for full activity.

In summary, BMPs 2, 3 and 7 are modified by phosphorylation (~33%) and glycosylation (50%). These post-translation modifications do affect protein morphogenic activity.

Matrix compositions useful in treating intervertebral disc impairment in vertebrates, including humans, may be prepared according to the foregoing descriptions and examples. While various embodiments of the inventions have been described in detail, modifications and adaptations of those embodiments will be apparent to those of skill in the art in view of the present disclosure. However, such modifications and adaptations are within the spirit and scope of the present inventions, as set forth in the following claims.

#### WHAT IS CLAIMED IS

- 1. A matrix for treating a patient having degenerative disc disease, the matrix comprising an injectable fluid comprising digestion-resistant remodelable collagen, said collagen being cross-linked through photooxidative catalysis and irradiation by visible light; and
- a plurality of living cells dispersed within said injectable fluid to form an injectable cell matrix for treating degenerative disc disease, said cells having inherent capability to elaborate proteoglycans in vivo.
  - 2. The injectable matrix of claim 1, further comprising
- a plurality of purified cell growth factors dispersed within said injectable cell matrix to form an injectable disc regeneration fluid, said living cells being responsive to said purified cell growth factors by increased elaboration of proteoglycans in vivo.
  - 3. The matrix of claim 1 wherein said cells are chondrocytes.
  - 4. The matrix of claim 1 wherein said cells are mesenchymal stem cells.
  - 5. The matrix of claim 4 wherein said cells are human-derived.
- 15 6. The matrix of claim 1 wherein said collagen is cross-linked using methylene blue as a photooxidative catalyst.
  - 7. The matrix of claim 1 wherein said cells are cultured in vitro to increase their response to said cell growth factors.
- 8. The injectable disc regeneration fluid of claim 2 wherein at least two of said plurality of cell growth factors are bone-derived.
  - 9. An injectable disc regeneration fluid, comprising

an injectable cell matrix according to claim 7; and

- a plurality of cell growth factors dispersed within said injectable cell matrix to form an injectable disc regeneration fluid, said living cells being responsive to said cell growth factors by increased elaboration of proteoglycans in vivo.
- 10. A method of treating a patient presenting with degenerative disc disease, the method comprising

providing an injectable disc regeneration fluid according to claim 2; and

injecting said injectable disc regeneration fluid into at least one of said patient's intervertebral discs to treat degenerative disc disease in said disc.

11. A method of continuing treatment of a patient presenting with degenerative disc disease, the method comprising

treating the patient according to the method of claim 10; and

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injecting a plurality of cell growth factors into said at least one of said patient's intervertebral discs after completion of the method of claim 10 to continue treatment of a patient presenting with degenerative disc disease.

- 12. A patient having a history of degenerative disc disease, wherein the patient has been treated by the method of claim 11.
  - 13. An intervertebral disc in vivo, said disc having been injected with injectable disc regeneration fluid according to claim 2.
  - 14. A method of treating a patient presenting with signs of hydrodynamic intervertebral disc dysfunction, the method comprising
- diagnosing hydrodynamic disc dysfunction in at least one intervertebral disc of said patient; testing said at least one intervertebral disc of said patient to establish cellular proteoglycan production within said at least one disc; and injecting a plurality of purified cell growth factors into said at least one disc to treat a patient presenting with signs of hydrodynamic intervertebral disc dysfunction.
- 15. The method of claim 14 wherein at least two of said plurality of cell growth factors are bonederived.
  - 16. An injectable cell growth medium for intervertebral disc regeneration, said medium comprising
  - an injectable fluid comprising digestion-resistant remodelable collagen, said collagen being crosslinked through photooxidative catalysis and irradiation by visible light; and
  - a plurality of purified cell growth factors dispersed within said fluid to form an injectable cell growth medium.
  - 17. The injectable cell growth medium of claim 16 wherein at least two of said plurality of cell growth factors are bone-derived.
- 25 18. An injectable disc regeneration fluid for intervertebral discs, the material comprising injectable cell growth medium according to claim 17; and cells responsive to said injectable cell growth medium through proteoglycan elaboration in vivo.
  - 19. The disc regeneration fluid of claim 18 wherein said cells are chondrocytes:
  - 20. The disc regeneration fluid of claim 18 wherein said cells are mesenchymal stem cells.
- 30 21. The disc regeneration fluid of claim 20 wherein said cells are human-derived.
- 22. The disc regeneration fluid of claim 20 wherein said cells are cultured in vitro to increase their response to said cell growth factors
  - 23. An injectable material for treating a patient for hydrodynamic disc dysfunction, the material made by a process comprising

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cross-linking collagen through photooxidative catalysis and irradiation by visible light; purifying a plurality of bone-derived cell growth factors;

dispersing said purified bone-derived cell growth factors within said cross-linked collagen; and dispersing cells responsive to said purified plurality of bone-derived cell growth factors within said cross-linked collagen to form an injectable material for treating hydrodynamic disc dysfunction.

- 24. The injectable material of claim 23 wherein said cells are chondrocytes.
- 25. The injectable material of claim 23 wherein said cells are mesenchymal stem cells.
- 26. The injectable material of claim 24 wherein said cells are human-derived.
- 10 27. The injectable material of claim 24 wherein said cells are cultured in vitro to increase their response to said cell growth factors.
  - 28. A method of hydrating an intervertebral disc annulus fibrosus in vivo, the method comprising

testing said disc for cellular proteoglycan production within said disc; and

- injecting cell growth medium according to claim 16 into said disc to hydrate the annulus fibrosus.
  - 29. A method of reducing susceptibility to herniation of an intervertebral disc in a patient having a history of intervertebral disc herniation, the method comprising testing said disc for cellular proteoglycan production within said disc; and injecting cell growth medium according to claim 16 into said disc to reduce susceptibility to herniation through hydration of the annulus fibrosus.
  - 30. A method of increasing the height of a patient presenting with hydrodynamic disc dysfunction in at least one intervertebral disc, the method comprising testing said at least one disc for cellular proteoglycan production within said at least one disc; and injecting cell growth medium according to claim 16 into said at least one disc to increase the height of said patient by increasing intervertebral spacing through increased proteoglycan production in said at least one disc.
  - 31. An injectable cell suspension for treating a patient having degenerative disc disease, the suspension comprising

an injectable fluid comprising a plurality of purified cell growth factors; and

- a plurality of living cells dispersed within said injectable fluid to form an injectable cell suspension for treating degenerative disc disease, said cells being responsive to said cell growth factors by increased elaboration of proteoglycans.
  - 32. The injectable cell suspension of claim 31 wherein said cells are chondrocytes.
  - 33. The injectable cell suspension of claim 31 wherein said cells are mesenchymal stem cells.

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- 34. The injectable cell suspension of claim 33 wherein said cells are human-derived.
- 35. The injectable cell suspension of claim 31 wherein said cells are cultured in vitro to increase their response to said cell growth factors.
- 36. The injectable cell suspension of claim 31 wherein at least two of said plurality of cell growth factors are bone-derived.
- 37. A method of treating a patient presenting with degenerative disc disease, the method comprising

providing an injectable cell growth medium according to claim 16; and

injecting said injectable cell growth medium into at least one of said patient's intervertebral discs to treat degenerative disc disease in said disc.

38. A method of treating a patient presenting with hydrodynamic disc dysfunction, the method comprising

providing an injectable material according to claim 23; and

injecting said injectable material into at least one of said patient's intervertebral discs to treat hydrodynamic disc dysfunction in said disc.

39. A method of treating a patient presenting with degenerative disc disease, the method comprising

providing an injectable cell suspension according to claim 31; and

injecting said injectable cell suspension into at least one of said patient's intervertebral discs to treat degenerative disc disease in said disc.

40. A method of cross-linking collagen to make digestion-resistant remodelable cross-linked collagen, the method comprising

providing a hydrogel comprising collagen;

solution for about 24 to about 72 hours.

containing said hydrogel within a semipermeable membrane, said membrane being substantially transparent to visible light and substantially permeable to at least one photooxidative catalyst;

transporting at least one photooxidative catalyst through said semipermeable membrane and into said hydrogel; and

irradiating said hydrogel with visible light to cross-link said collagen.

- 41. The method of claim 40 comprising an additional step between said containing step and said transporting step, the additional step being submerging said hydrogel-containing semipermeable membrane in a high salt:high sucrose
  - 42. The method of claim 41 wherein said hydrogel comprises nucleus pulposus tissue.

- 43. The method of claim 42 wherein said transporting step occurs at a substantially constant hydrogel temperature of about 10°C.
- 44. Digestion-resistant remodelable cross-linked collagen made by the method of claim 43.
- 45. The method of claim 41 wherein said hydrogel comprises substantially Type II collagen.
- 46. The method of claim 43 wherein said semipermeable membrane comprises dialysis tubing having a molecular weight cutoff of about 3500 Daltons.
  - 47. The method of claim 43 wherein said transporting step and said irradiating step are substantially simultaneous.
- 48. The method of claim 43 wherein said transporting step occurs while said semipermeable membrane containing said hydrogel is immersed in a solution comprising at least one photooxidative catalyst.
  - 49. The method of claim 43 comprising an additional step after said irradiating step, the additional step being
- terminating said transporting step and said irradiating step when said collagen cross-linking is substantially complete.
  - 50. The method of claim 49 comprising an additional step after said terminating step, the additional step being

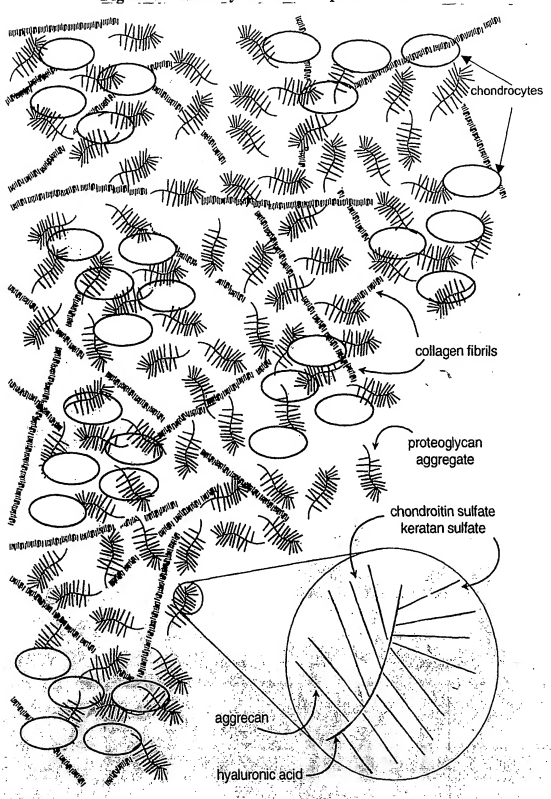
extracting said cross-linked collagen from said hydrogel.

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- 51. The method of claim 43 wherein said at least one photooxidative catalyst comprises methylene blue.
  - 52. A fluid matrix for treating intervertebral disc disease in a vertebrate, said fluid comprising nucleus pulposus tissue of a donor vertebrate.
  - 53. The fluid matrix of claim 52 wherein said nucleus pulposus tissue is cross-linked.
  - 54. The fluid matrix of claim 53 further comprising a growth factor.
- 25 55. The fluid matrix of claim 54 further comprising a plurality of living cells.
  - 56. The fluid matrix of claim 55, wherein said plurality of living cells comprise chondrocytes.
  - 57. The fluid matrix of claim 55, wherein said plurality of living cells comprise mesenchymal stem cells.
  - 58. The fluid matrix of claim 55, wherein said plurality of living cells are human-derived.
- 30 59. The fluid matrix of claim 52, wherein said nucleus pulposus tissues are decellularized.

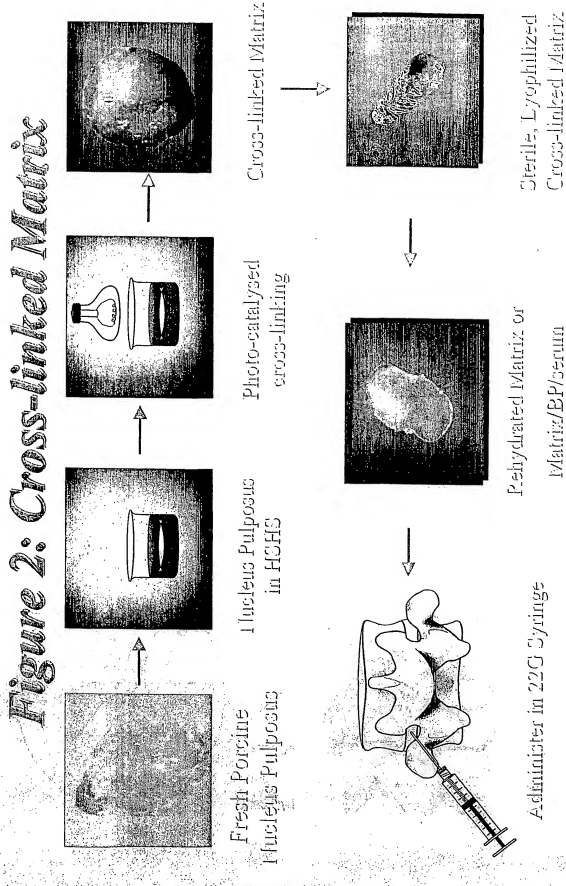
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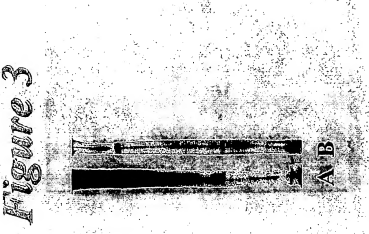
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Figure 1 – Healthy Nucleus Pulposus Tissue



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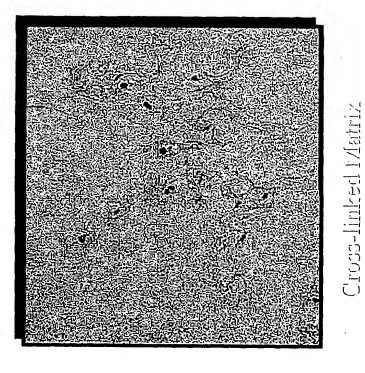
ne A. Mon cross-linked control shows substantial protein extraction by E. Cross-linked matrix demonstrates reduced protein extraction

disrupted, crenated cell fragments

minimal cell membrane material

further isopropanol Sterilization

# Incloses Pulposus versus Cross-linked Matrix rigure 4: Comparison of Fresh Porcine

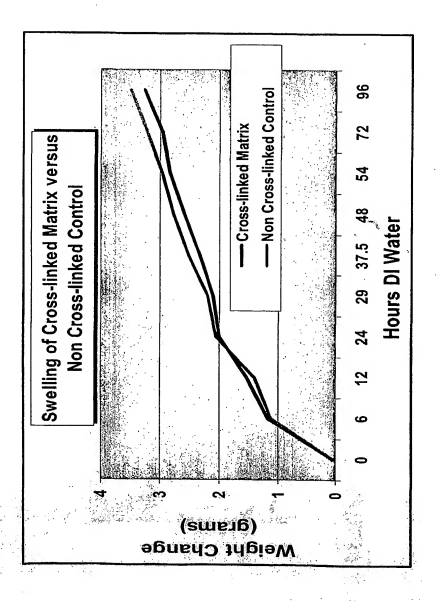


Fresh Mucleus Pulposus

• intact pericellular matriz "nests" round, nucleated chondrocytes

Sang A Peppin digests of non cross-linked control react with Type II collagen antibodies

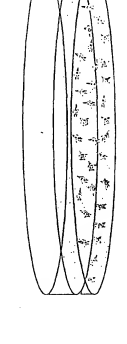




-Cross-linked matrix retains 95% hydraulic capacity

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# Growth and Proliferation of Disc cells into Cross-linked Matrix Cell Isolation Culture Expansion

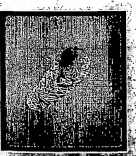


Cross-linked Matrix/EP Disc Cell Culture in



Cross-linked matriz/EP



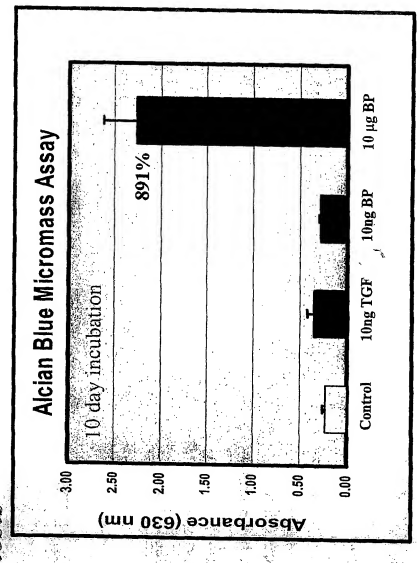


Shoss-lifited, mairiz

enzymatic digestion for pulposus from sheep Sterile nucleus spine;

cell isolation

Figure 8: Growth Factor Stimulation of Matrix

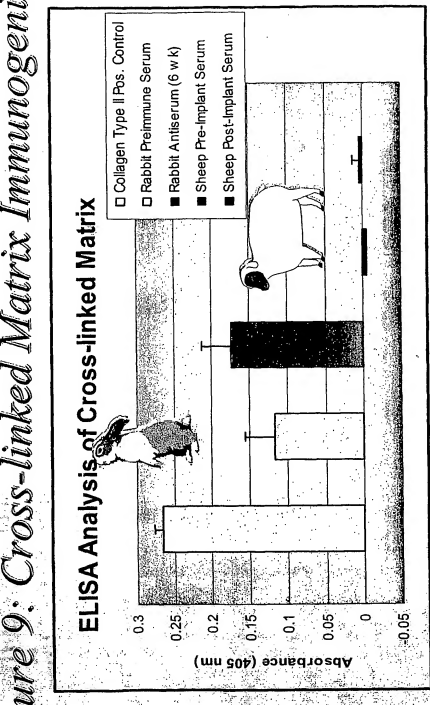


Significant stimulation of matrix production only at 9 g EP concentrations

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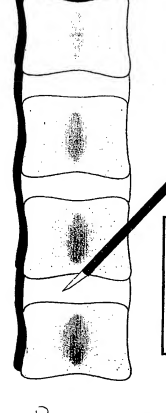


initionizations

alle serum antibodies to cross-linked matrix *in vivo* (1st sheep)



Stoperation: annulus stabs to create two degenerative discs



Wait 2 months

Znd operation: Cross-linked

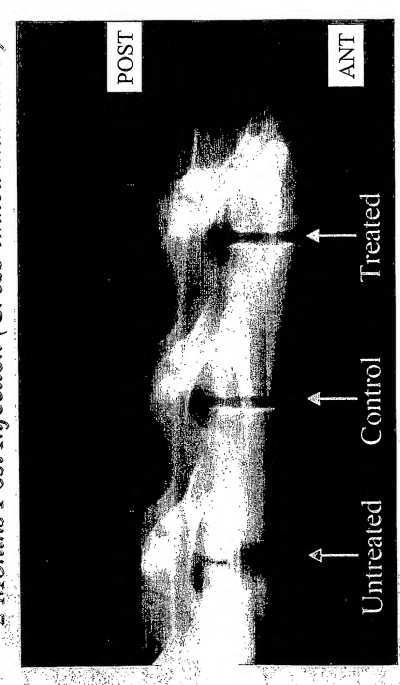


one disc

Sacrifice: 2, 4, and 6 months

Histomorphometry MRV radiographs

Figure II: Radiograph-Pilot Study #1 2 Months Post Injection (Cross-linked matrix/BP)



- Treated and Control discs; normal size and appearance of disc

Structure

· Untreated disc: disjunct endplates, bone resorption and remodeling

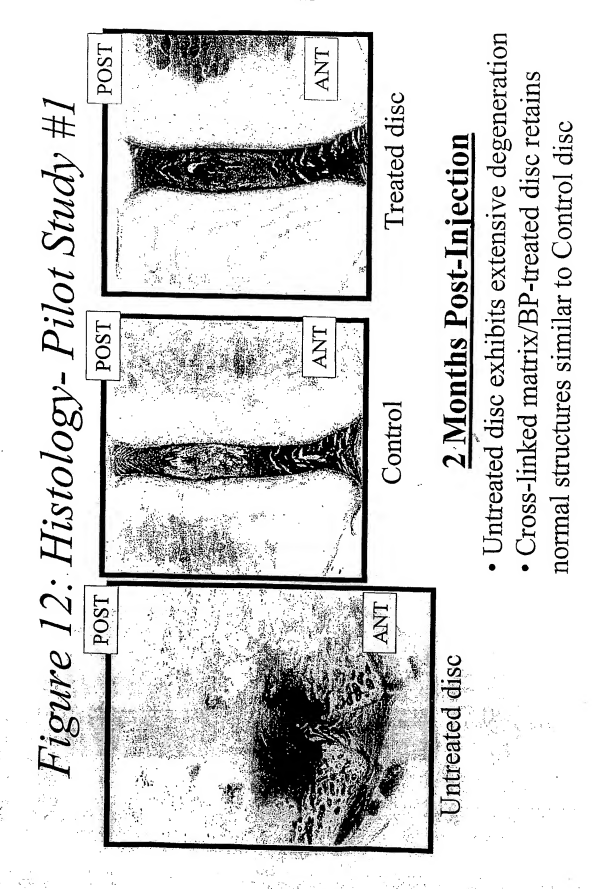
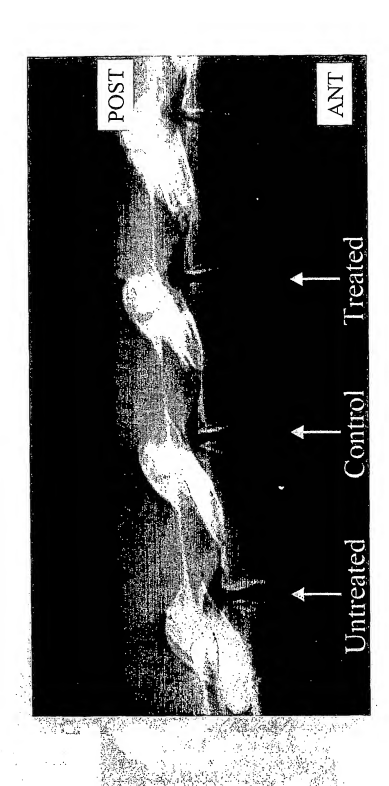
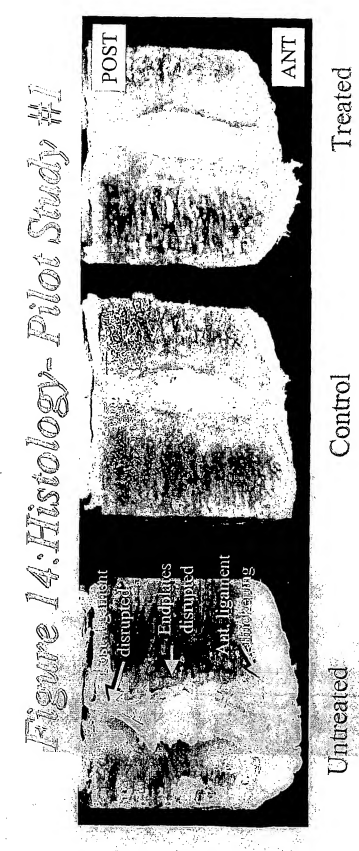


Figure 13: Radiograph Pilot Study #1 4 Months Post Injection (Cross-linked matrix/BP)



110 apparent radiographic differences between discs in 4 month sheep



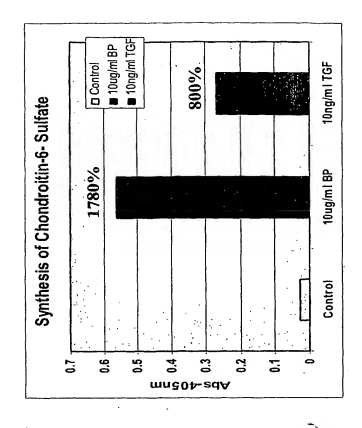
Treated

4 Months Post-Injection

S. Cross-linked matrix/BP-freated disc similar to control disc. normal gelatinous nucleus, regular annulus, intact endplates Well intreated disc exhibits degenerative changes

Intreated

Figure 15: Growth Factor Stimulation of If Collagen & Chondroitin-6-Sulfate Synthesi.



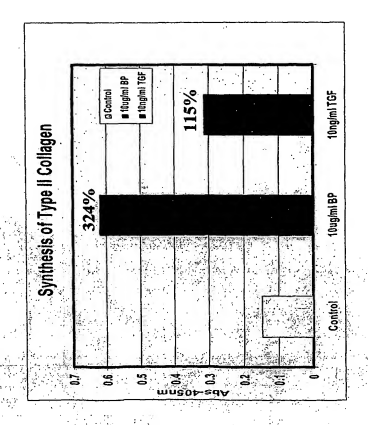
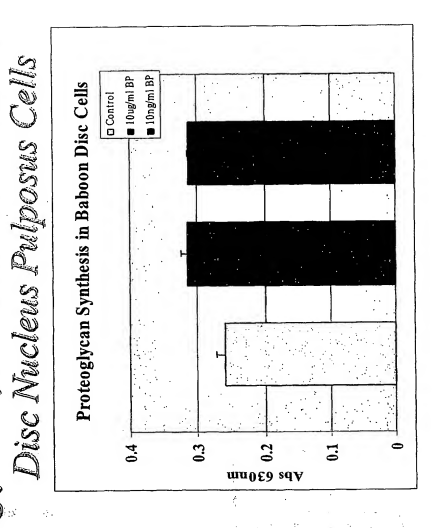


Figure 16: Growth Factor Stimulation of Proteoglycan

Synthesis in Fuman Intervertebral Disc Nucleus Pulposus Cells ■ 10ug/ml BP ■ 10ng/ml TGF Control Proteoglycan Synthesis in Human Disc Cells Figure 16b 9 day incubation 9.0 0.4 0.7 Mn0£9 sdA ■ 10mg/ml TGF ■ 10ug/ml BP ■ 20ug/ml BP Proteoglycan Synthesis in Human Disc Cells Figure 16a 8 day incubation

oteoglycan Synthesis in Baboon Intervertebra sigure 17: Growth Factor Stimulation of



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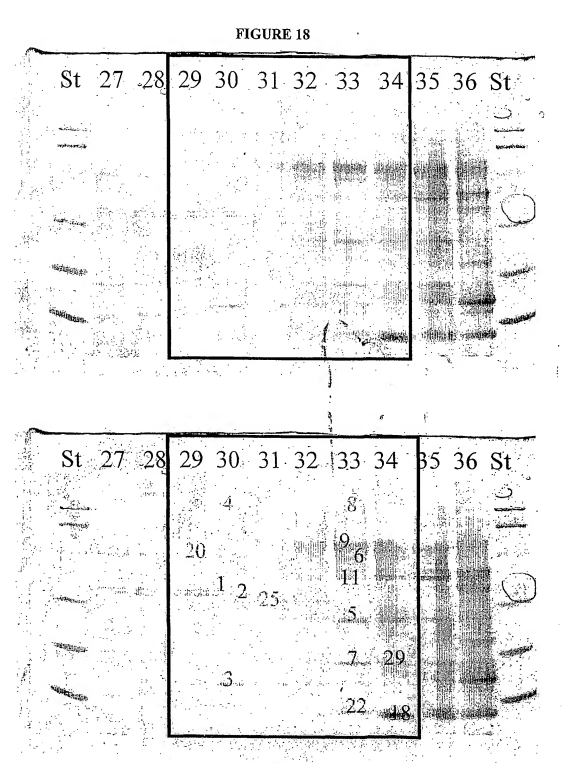
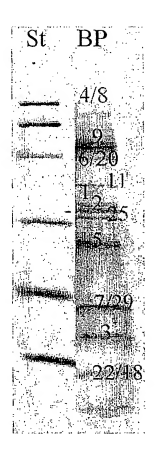


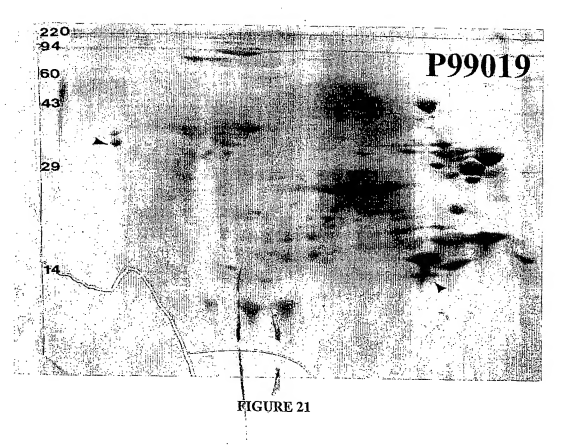
FIGURE 19



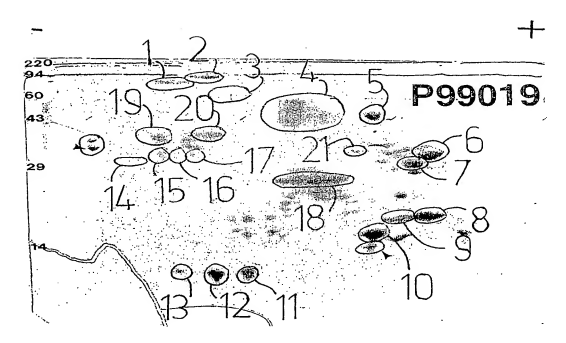
Band No.	Identity	
1	Histone H1.c	
2	Histone H1.c	
3	Ribosomal protein RS20	
4	Similar to ribosomal protein LORP	
5	BMP-3	
6	α2 macroglobulin RAP and BMP-3	
7	Similar to ribosomal protein LORP	
8	BMP-3	
9	BMP-3	
11	Ribosomal protein RL6 and BMP-3	
18	TGF-β2 / SPP 24	
20 .	Factor H	
22	TGF-β2	
25	BMP-3 and H1.x	
29 .	BMP-3 and ribosomal protein RL32	

FIGURE 20

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No.	Identity	No.	Identity
1	Factor XIII	11	TGF- β2/SPP24
2	LORP	12	SPP24
3	LORP	12	TGF- β2/SPP24
4		14	lysyl oxidasc
5	RL3	15	lysyl oxidase
6		16	lysyl oxidase
7		17	lysyl oxidase
8		18	BMP-3
9	-	19	cathepsin L
10		20	
		21	RŠ3a

FIGURE 22

Timed Ion Selector: 16.1 OFF

Pressure: 1.70e-06

Scans Averaged: 256

Grid Voltage: 94.000 %

Accelerating Voltage: 20000

Mode: Linear

Guide Wire Voltage: 0.075 %

Delay: 50 ON

Laser: 1965

Low Mass Gate: 500.0

PSD Mirror Ratio:

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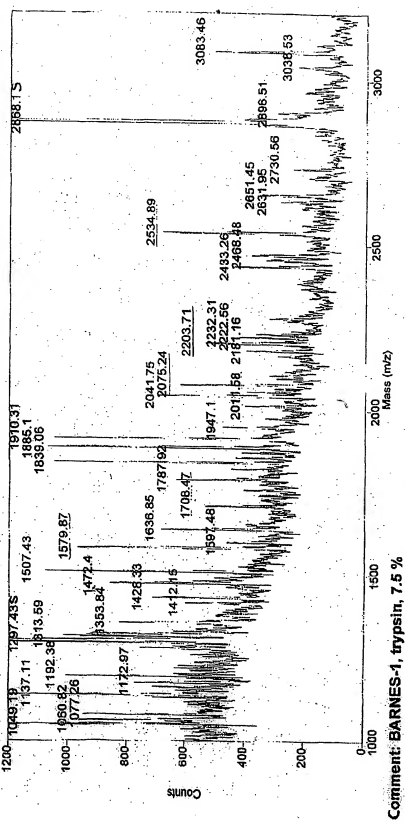
Mirror Ratio: 1.060

Figure 23A (Band 1)

## Core Protein ( Columbia University /HHM

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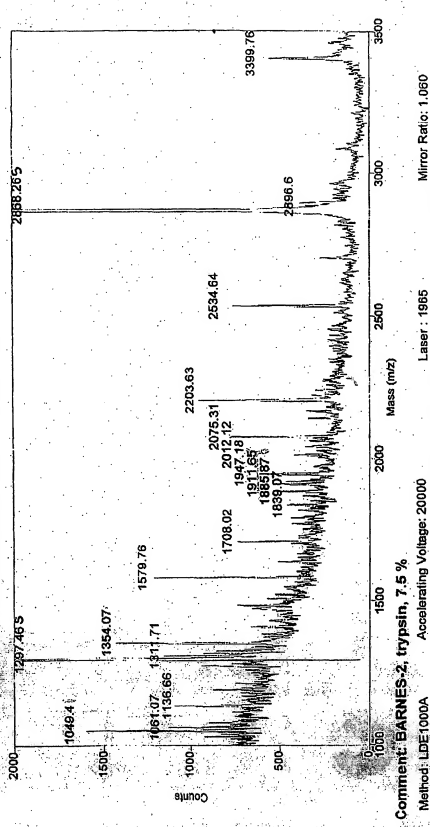
Negative Ions: OFF

Figure 23B (Band 2)

## Protein Core Columbia University /HH

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Negative lons: OFF

Jow Mass Gate: 500.0

Delay: 50 ON

Figure 23C (Band 3)

## 3876.36 Sample: 76 Savitsky-Golay Order = 2 Points = 19 Timed Ion Selector: 16.1 OFF Mirror Ratio: 1.060 Protein Core PSD Mirror Ratio: Collected: 10/12/99 2:31 PM 3404.02 2979.08 2868.08 S 30,00 Pressure: 1.5' e-06 Laser: 1965 Scans Averaged: 256 Columbia University /HHMI This File # 1: C: NOYAGERIDATAIMAG1099IDIGESTISMOOTH.MS Original Filename: c:\woyager\data\mag1099\digest\bare028.ms Mass (m/z) 2307.55 Grid Voltage: 94.000 % Guide Wire Voltage: 0.075 % Accelerating Voltage: 20000 2002 1919.02 1649.68 Comment: BARNES-3, trypsin, 7.5 % 1500 Method: LDE1000A Sounts

Timed Ion Selector: 16.1 OFF

Pressure: 9.04e-07

Guide Wire Voltage: 0.075 %

Low Mass Gate: 500.0

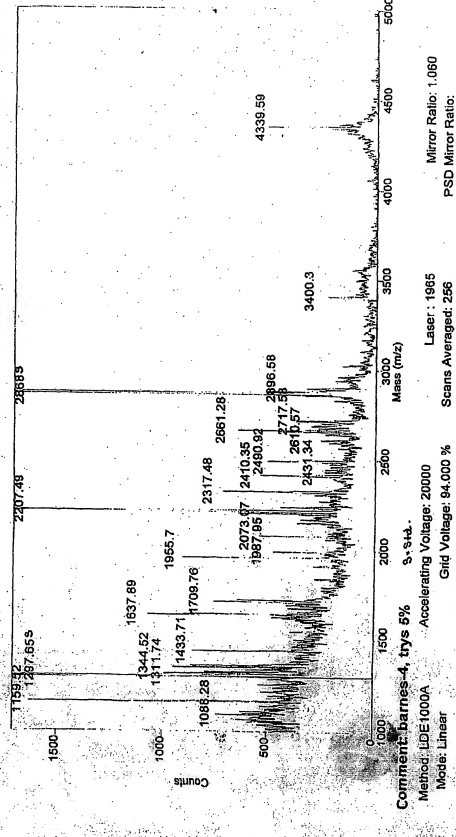
Negative lons: OFF

Figure 23D (Band 4)

## **Protein Core** University /HHM Columbia |

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Savitsky-Golay Order = 2 Points = 19 Collected: 10/27/99 2:30 PM Sample: 22



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Pressure: 3.68e-07

Scans Averaged: 121

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Guide Wire Voltage: 0.075 %

Delay: 50 ON

Low Mass Gate: 500.0

Laser: 1965

PSD Mirror Ratio:

Negative lons: OFF

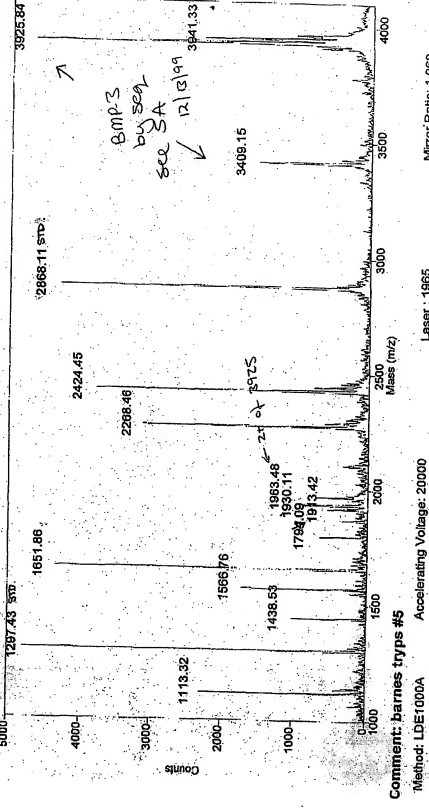
Mirror Ratio: 1.060

Figure 23E (Band 5)

## **Protein Core** Columbia University /HHM

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Timed Ion Selector: 16.1 OFF

Pressure: 4.06e-07

Scans Averaged: 256

Grid Voltage: 94.000 %

Accelerating Voltage: 20000

Method: LDE1000A

Guide Wire Voltage: 0,075 %

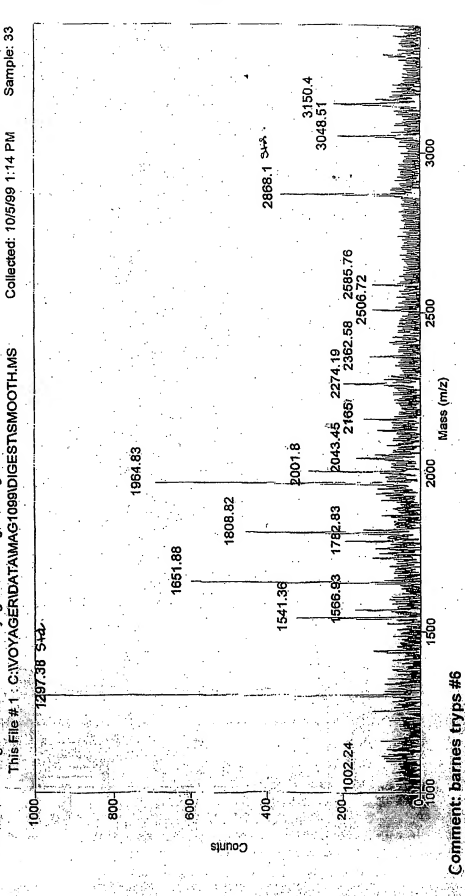
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PSD Mirror Ratio:

Mirror Ratio: 1.060





Timed Ion Selector: 16 1 OFF

Pressure: 5.89e-07

Guide Wire Vollage: 0.075 %

Delay: 50 ON

Low Mass Gate: 500.0

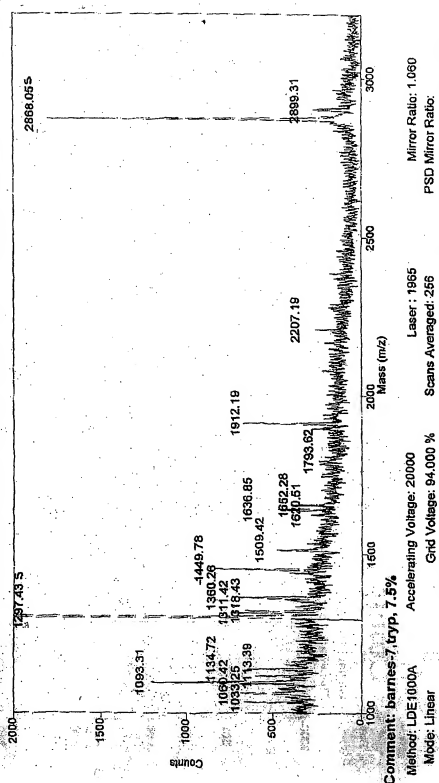
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Figure 23G (Band-7)

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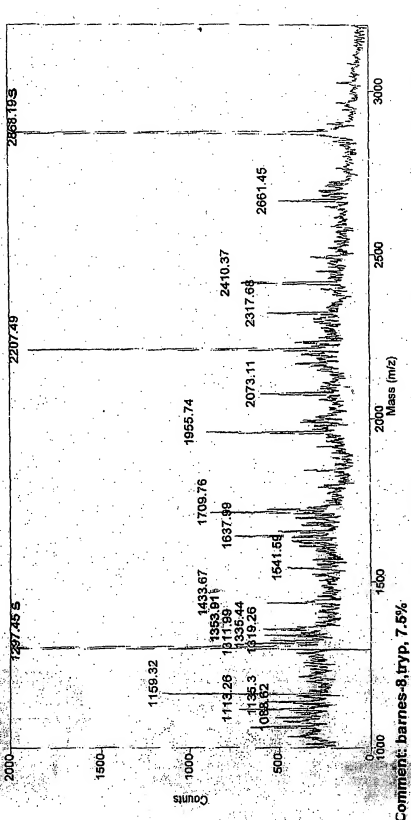
Counts

Figure 23H (Band 8)

# Protein Core

IE# 1:: C:IVOYAGERIDATAIMAG1199IDIGESTISMOOTH.MS Original Filename: c:\voyager\data\mag1199\digest\snow\_005.ms

Sample: 64 Savitsky-Golay Order = 2 Points = 19 Collected: 11/10/99 3:18 PM



Accelerating Voltage: 20000

Grid Voltage: 94.000 % Guide Wire Voltage; 0.075 %

Delay: 50 ON

Laser: 1965 Scans Averaged: 256

Pressure: 4.01e-07 Low Mass Gate: 500.0

Mirror Ratio: 1.060 PSD Mirror Ratio:

Timed Ion Selector: 16.1 OFF Negative Ions: OFF



Protein Core Columbia University /HHMI

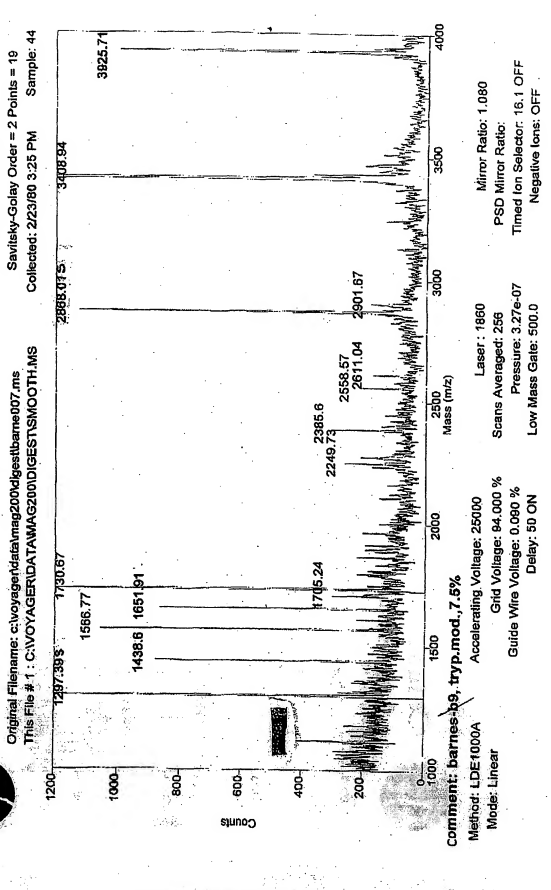
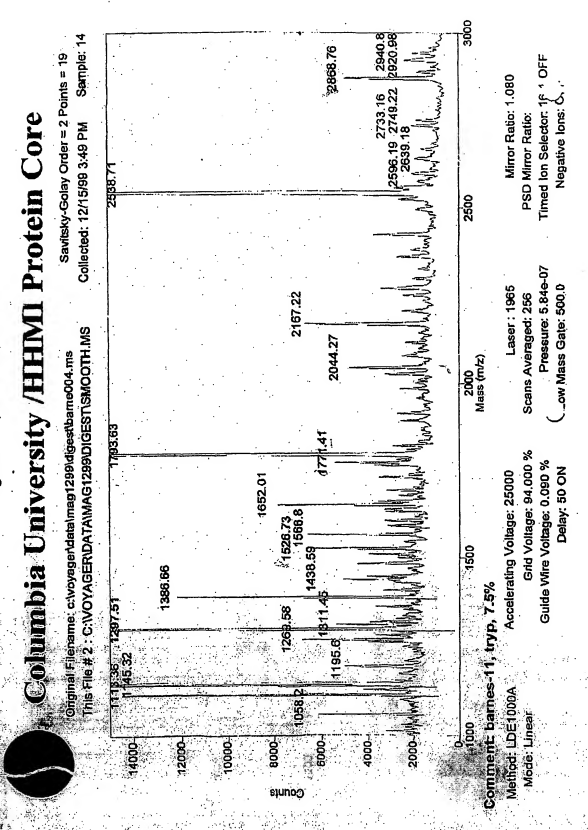


Figure 23J (Band 11)



Timed Ion Selector, 16.1 DFF

Pressure: 2.90e-07

Scans Averaged: 258

Grid Voltage: 94.000 %

Guide Wire Voltage: 0.090 %

Delay: 50 ON

Low Mass Gate: 500.0

Laser: 1745

PSD Mirror Ratio:

Negative lons: OFF

Mirror Ratio: 1.080

Figure 23K (Band 18)

## **Protein Core** umbia University /HHM

his File # 2: C:IVOYAGERIDATAIWAG12991DIGESTISMOOTH.MS Original Filename: c:\voyager\data\mag1299\digest\bame005.ms

Sample: 13 Savitsky-Golay Order = 2 Points = 19 Collected: 12/15/99 4:47 PM

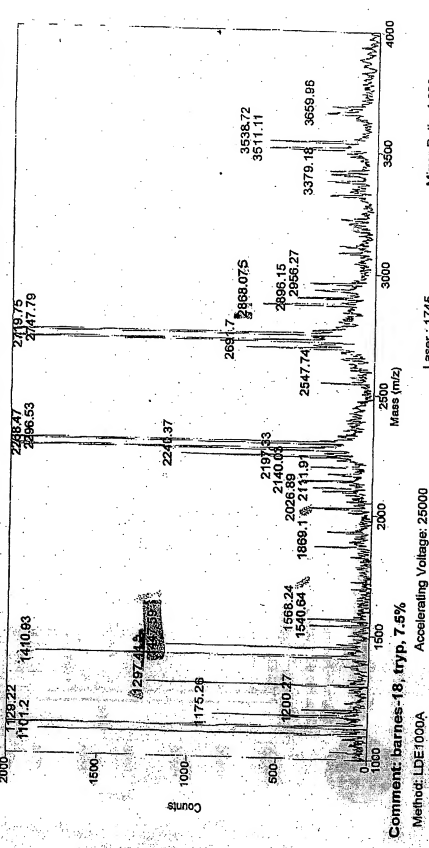


Figure 23L (Band 20)

## **Protein Core** Columbia University /HHN

This File # 4 C. WOYAGERIDATAIMAG10001GESTISMOOTH.MS Original Filename: c:\voyager\data\mag100\digest\bame001.ms

2073.84

Savitsky-Golay Order = 2 Points = 19 Collected: 1/6/80 3:36 PM

Mirror Ratio: 1.080 Negative lons: Or Timed Ion Selector: PSD Mirror Ratio: يموريهم أيواك أولي بدريه ويهومهم والمعادية المالية المالية المالية المالية

Mass (m/z)

2500

Laser: 1820 Scans Averaged: 256

Grid Voltage: 94.000 %

Accelerating Voltage: 25000

Guide Wire Voltage: 0.090 %

Delay: 50 ON

Low Mass Gate: 500.0

Pressure: 9.21e-07

2202.07

1802.96

Figure 23M (Band 22)

## **Protein Core** University /HHM

This File # 1: C: WOYAGERIDATANMAG2001DIGESTISMOOTH.MS Original Filename: c:\voyager\data\mag200\digest\barne003.ms

Collected: 2/16/80 3:35 PM

Sample: 54

Savitsky-Golay Order = 2 Points = 19

2867.945 9.68 2547.71 2384.16 2240.25

1838.16

1664.67

Laser: 1870 Mass (m/z)

2500

...ow Mass Gate: 500.0 Scans Averaged: 231

**Grid Voltage: 94.000 %** 

Guide Wire Voltage: 0.090 %

Delay: 50 ON

Accelerating Voltage: 25000

mod.tryp., 7.5%

Pressure: 3.43e-07

Mirror Ratio: 1.080 PSD Mirror Ratio:

Timed Ion Selector: 16 1 OFF Negative lons: 🖒

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Counts

Timed Ion Selector: 16 'OFF

Negative lons: Orr

Low Mass Gate: 500.0

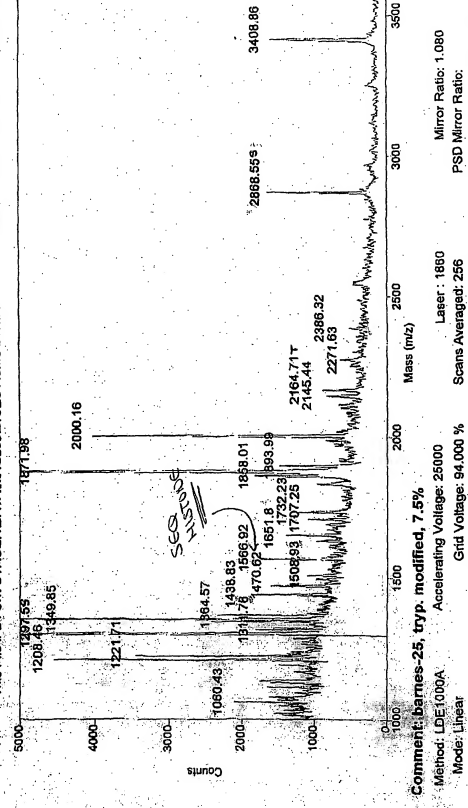
Guide Wire Voltage: 0.090 %

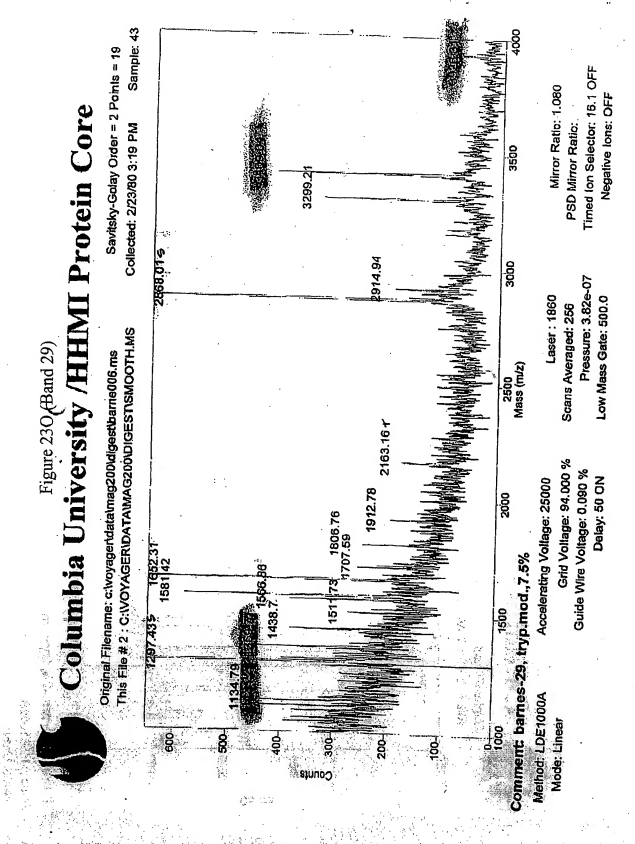
Delay: 50 ON

Figure 23N (Band 25)

## Protein Core Columbia University /HHM

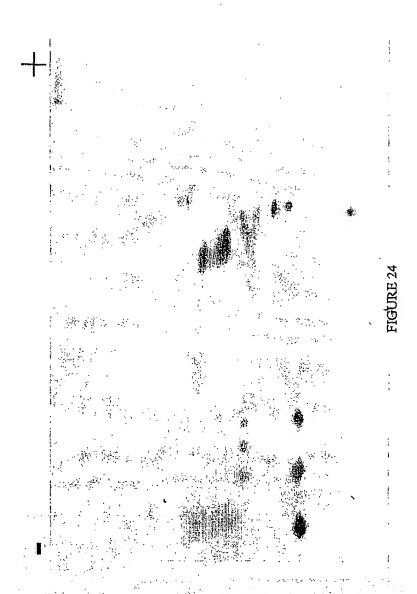
Savitsky-Golay Order = 2 Points = 19 Collected: 2/2/80 3:24 PM This File # 2: C:NOYAGERIDATAMAG200DIGESTISMOOTH.MS Original Filename: c:\voyager\data\mag200\diges\\\bane001.ms





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BNSDOCID: <WO 0176654A1 1 3



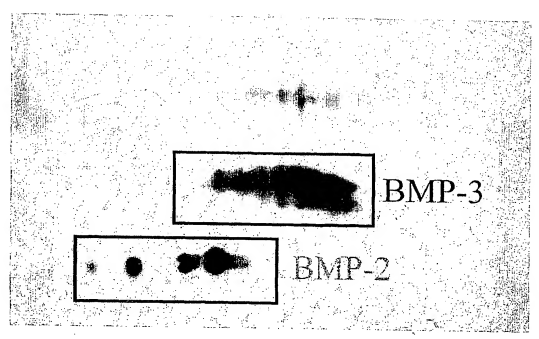


FIGURE 25A

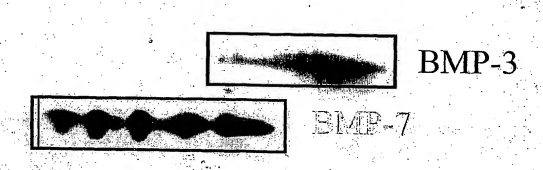


FIGURE 25B

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BNSDOCID: <WO 0176654A1 I

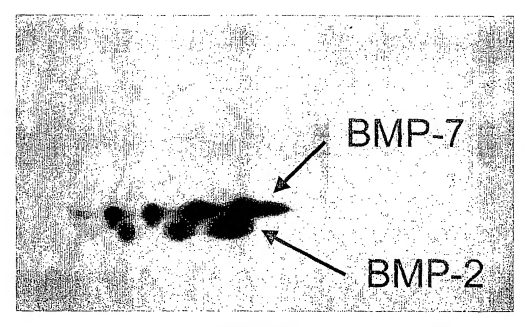


FIGURE 25C

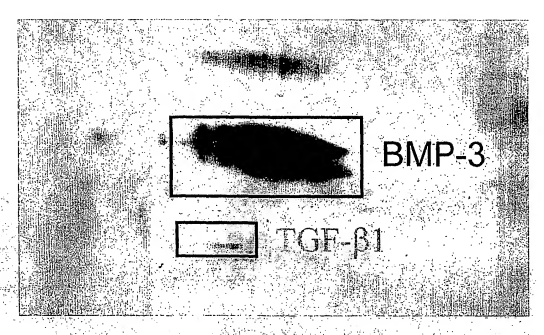
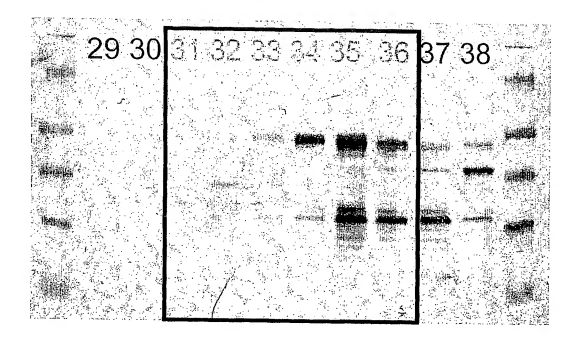
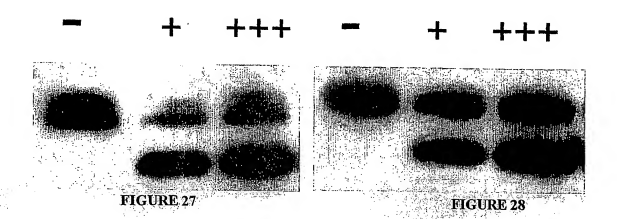


FIGURE 25D

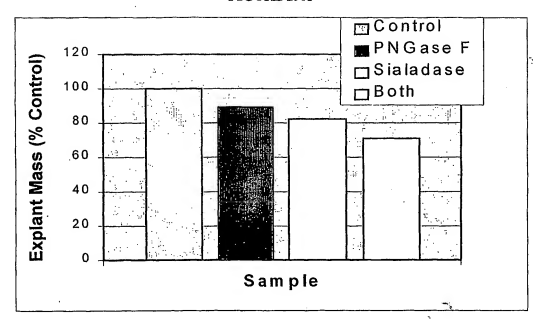
## FIGURE 26



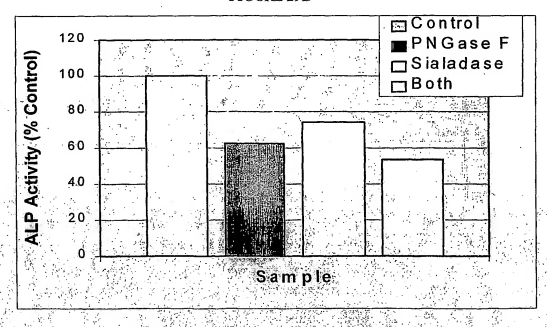


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FIGURE 29A



#### FIGURE 29B



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FIGURE 30
Antibody Information

Specificity	Antigen	Host Species	PC/MC	Source	Catalog No.
TGF-β1 (human)	Protein	Rabbit	Polyclonal	Promega	G1221
TGF-β2 (human)	Peptide	Rabbit	Polyclonal	Santa Cruz Biotechnology	sc-90
TGF-β3 (human)	Peptide	Rabbit	Polyclonal	Santa Cruz Biotechnology	sc-82
BMP-2 (human)	Protein	Rabbit	Polyclonal	Austral Biologics	PA-513-9
BMP-3 (human)	Peptide	Chicken	Polyclonal	Research Genetics	NA NA
BMP-4 (human)	Peptide	Goat	Polyclonal	Santa Cruz Biotechnology	sc-6896
BMP-5 (human)	Peptide	Goat	Polyclonal	Santa Cruz Biotechnology	sc-7405
BMP-6 (human)	Peptide	Mouse	Monoclona l	Novocastra Laboratories	NCL-BMP6
BMP-7 (human)	Peptide	Rabbit	Polyclonal	Research Genetics	NA
FGF-1 (human)	Peptide	Goat	Polyclonal	Santa Cruz Biotechnology	sc-1884
osteonectin	Protein	Mouse	Monoclona	DSHB	AON-1
(bovine) osteocalcin (bovine)	Protein	Rabbit	Polyclonal	Accurate Chemicals	A761/R1H
serum albumin (bovine)	Protein	Rabbit	Polyclonal	Chemicon International	AB870
transferrin (human)	Protein	Chicken	Polyclonal	Chemicon International	AB797
apo-A1 lipoprotein (human)	Protein	Goat	Polyclonal	Chemicon International	AB740

Figure 31A. Identification of Proteins by Amino Acid Sequencing of Tryptic Fragments

			407-84	Andon	- Acatification	Spariae	Arc No.	AAs
Band	Band Sample	Sequence Data	Best Darapase Marchiwarch Incentification	Match	Identification	T	200	
						·		
2	fx 49	XLAAAGYDVEK	ALAAAGYDVEK	11/11	histone H1.c	human	87668 (NCBI)	65-75
m.	(8/61)	SLEKVCADLIR	SLEKVCADLIR	11/11	40s Ribosomal Protein	rat	R3RT20 (PIR)	31-41
4	(1340) fx 65 ()	(V)VCGMLGFPSEAPV	GFPSEAPV VVCGMLGFPGEKRV 11/14	11/14	LORP	mouse	AAC95338 (NCBI)	213- 226
5	N terminal	STGVLLPLONNELPG	STGVLLPLQNNELPG 15/15	15/15	ВМР-3	human	4557371 (NCBI)	290- 304
And the second second	(x 72	STGVLLPLONNELPGA FYOY	STGVLLPLONNELPG 20/20 AEYOY	20/20	BMP-3	human	4557371 (NCBI)	290- 309
<b>.</b>	fx 74	STGVLLPLQ	PLQ	6/6	BMP-3	human	4557371 (NCBI)	290- 298
ပ	(575) (4.566)	(s)атгағхе	SQTLQFDE	8/2	BMP-3	human	4557371 (NCBI)	346- 353
	7.P.34	WAF	no match.		222			
	N terminal	HAGKYSREKNT(P)A(P	HGGKYSREKNQPKP	11/14	α2-Macroglobulin Receptor Assoc. Pro.	исшаи	P30533 (Swiss-Prot)	31-46
	fx 57	SQTLQFDEQ	SQTLOFDEQ	6/6	BMP-3	human	4557371 (NCBI)	346- 354
	fx 57 (1852)	SLKPSNHA	SLKPSNHA	8/8	BMP-3	human		410- 417
7	fx 51	AALRPLVKP	AALRPLVKP	6/6	60s Ribosomal Protein L32	esnow	P17932 (Swiss-Prot)	1-9
	1×37 (no MS)	A(H))(Q)VERYV	AIVER	5/2	60s Ribosomal Protein L32	mouse	rot)	109- 113
	fx 37 (no MS)	A(H)I(Q)VERYV	наѕркүу	2/2	60s Ribosomal Protein L32	mouse	P17932 (Swiss-Prot)	22-28
8	fx 78 0	XALF(G)AQLGXALGPI	no match		155			
	fx 56 (1587)		DEQT	10/10	BMP-3	human	P12645 (Swiss-Prot)	346- 355
	,,,,,,							

Identification of Proteins by Amino Acid Sequencing of Tryptic Fragments

Band	Band  Sample	Seguence Data	Best Database Match Match		Identification	Species	Acc. No.	AAs
E	fx 55		SQTLQF	5/6	BMP-3	human	4557371	346-
	(1311)						(NCBI)	351
. 21	fx 47	VLATVTKPVGGDK	VLATVTKPVGGDK	13/13	60s Ribosomal Protein L6	human	Q02878	87-99
	(1772)		-				(Swiss-Prot)	
1.1	92 ×3	XVFAL	VFAL	4/4	60s Ribosomal Protein L6	human	Q02878	273-
	(1795)						(Swiss-Prot)	276
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	fx 61	AVPOLOGYLR	AIPOLOGYLR	9/10	60s Ribosomal Protein L6	human	Q02878	262-
-0.50	(1145)		*				(Swiss-Prot)	271
						: : :		
- 8 <u>-</u>		が、 は、						
22	fx 58	ALDAAYCFR	ALDAAYCFR	6/6	TGF-82	human	P08112	303-
- N	(1101)						(Swiss-Prot)	311
A. 40	fx 69 (no	GYNANFCAGACPYL	GYNANFCAGACPYL 14/14	14/14	TGF-82	human	P08112	340-
n koja vedem							(Swiss-Prot)	353
	fx 66	VNSQSLSPY	VNSQSLSPY	6/6	SPP24	bovine	Q27967	42-50
	(1411.71)						(Swiss-Prot)	
22	6£ x3	KAAKPSV(P)	KAAKPSVP	8/8	Histone H1.x	human	JC4928 (PIR)	199-
	(1470)							50g
29								
fx = fr	ction numb	fx = fraction number (molecular weight of	ar weight of fragment, as measured by SDS-PAGE)	by SD	S-PAGE)			

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e32A. Identification of Proteins by Mass Spectrometry of Tryptic Fragments

Comments			15 MS peaks match with					identification of starred	sequence analysis		15 MS peaks match with	Dallo										12 MS peaks match with Band 8			
% Cover-	age		<b>5</b> 5	:	-	• •		16		• ;			.62								·	∞			
AAc	}		110-121		65-79	64-79	35-54	65-79*		64-79	35-54		20-59	•:		76-83	56-66	88-99	9-21	5-21	88-119	150-167		648-669	455-478
Macc Diff.	erence		0.60		0.16	85.0	-0.74	0.05		0.13	-0.20		0.36			-0.09	-0.16	0.55	0.27	-0.17	-0.85	7 -0.32		-0.28	0.47
14000	Spec	Database	1172.37		1579.71	1707.89	2012.32	1579.71		1707.89	2012.32		1129.40			1156.30	1334.62	1351.58	1517.77	1919.19	3404.87	1988.27		2410.63	2610.10
	Spec	Data	1172.97		1579.87	1708.47	2011,58	1579.76		1708.02	2012.12		1129.76	. · 		1156.21	1334.46	1352.13	1518.04	1919.02	3404.02	1987.95		2410.35	2610.57
	Acc. No.		87668	(NCBI)				87668	(NCBI)	**:			R3RT20	(PIR)		•						NP002309	Prot)		
	Species		human					human			· · . · . : · :		rat									human			
	Mass Spec Profile		4 peaks	match with				3 peaks	match with			1	7 peaks	match with	ribosome	2 × × × × × × × × × × × × × × × × × × ×				ではない		3 peaks	Lysyl	Oxidase Rr	
1.5	Band		-					2				100	3							1		4			

32.B. Identification of Proteins by Mass Spectrometry of Tryptic Fragments

	% Cover- Comments	age	48 % coverage calculation is	3, 183 AAS (290-472)						identification of starred	peptide confirmed by sectionce analysis		17						15 % coverage calculation is	relative to the mature BMP-3, 183 AAS (290-472)		
	AAs %(		361-368		346-357	345-357	410-424	346-360	374-392	373-392		290-318*	283-290				129-150	257-282	346-357		410-424	
	Mass Diff-	erence	10.01		3 -0.05	00:0	0.05	0.07	3 -0.17	96.0-		1.38	60:00				0.15		0.18	*	-0.03	
		Spec Database	1113.31		3 1438.58	6 1566.76	6 1651.91	9 1794.02	6 2268.63	5 2424.81		5 3407.77	4 1002.15				8 2362.43	1 3048.52	1566.75		3 1651.91	
		Spec Data	71 1113.32	<u> </u>	1438.53	1566.76	1651.86	1794.09	2268.46	2424.45		3409.15	13 1002.24	<u>.</u>	· · · · · · · · · · · · · · · · · · ·		2362.58	3048.51	1566.93		1651.88	
	es Acc. No								<u> </u>				n P30533	Swiss-	Prot)				-	(NCBI)		
	ec Species		human	<b>.</b>		ág.		. 0005 240			u par		human	<u>-</u>		<b>.</b>			human	di a	3	:
A Company of the State of	nd Mass Spec	Profile	**************************************	march With BMP-3				777					3 peaks	matchwith	28	Macroglobuli n RAP			2 peaks	C20: 1		
Sec.	Band	andre	•	ဂ	Az er ni de				varios e				9			Ζ11. 15 (A.)		i e .			Ž.	

re32.C. Identification of Proteins by Mass Spectrometry of Tryptic Fragments

				•	•		•																	
Comments									% coverage calculation is	relative to the mature BMP- 3 183 AAS (290-472)					12 MS peaks match with	Dana 4		% coverage calculation is	relative to the mature BMP-3 183 AAS (290-472)					
% Cover-	age	,	33				:		21			:			က			36						
AAs	· ·		67-75			1-10*	65-74	19-29	102-111		361-368	190-200	410-424	346-360	648-669	· .		361-368		346-357		410-424		290-318
Mass Diff-	erence		0.08			-0.09	0.44	0.12	0.22		0.08	-0.32	0.37	-0.40	-0.26			-0.17		0.05	0.01	0.30	0.48	1.17
Mass	Spec	Database	1033.17		.* . 	1093.40	1134.28	1449.66	1060.20		111331	1360.58	1651.91	1794.02	2410.63	••		1113 31		1438.58	1586.76	1651.61	2901.19	3407.77
Mass	Spec	Data	1033.25			1093.31	1134.72	1449.78	1060.42		1113 30	1360.26	1652.28	1793.62	2410.37			1112.14	3	1438.60	1566.77	1651.91	2901.67	3408.94
Acc No			P17932	(Swiss-	Prot)			. :	4557371	(NCBI)	•			: :	NP002309	(Swiss-	Prot)	AEE 7274	(NCBI)	. •				
Chorine	Shecies		mouse				•		human						human	•		domina	in in in			: ".		
	) U		4 peaks	match with	ribosome	707			5 peaks	match with	5-HWH-3			ないというという	1 peak	matches with	Lysyl Oxidase RP	27000	match with	BMP-3				,
-	pand		17				S .								8	12.			D		71. A.			

e32.0. Identification of Proteins by Mass Spectrometry of Tryptic Fragments

_			_													_									_	_	
Commonto		% coverage calculation is	relative to the mature BMP-	3, 183 AAS (290-472)													*										
201100 /0	age	48							16			2.					52	÷.	•			30		•			
200	Ş	361-368			410-424	346-360	373-392	290-318	114-122		141-155	10.20	03-01		262-271	260-271	303-311		400-409	312-328	340-362	42-53		113-124	86-98	62-77	33-53
Maco Diff	erence	-0.08	٠.		-0.18	-0.44	75.0-	0.57	0.15		0.02	0.03	9		0.01	0.06	90.0-		-0.16	-0.23	-0.21	-0.67		-0.06	0.04	0.05	-0.10
Maco	Spec Database	1113.31			1651.91	1794.02	2424.81	3407.77	1140.23		1526 86	1050 12	1000		1145.35	1386.68	1101.26		1175.42	2240.60	2691,91	1411.60	· R	1447.65	1540.60	1869.05	2268.57
Mann	Spec Data	1113.23			1651.73	1793.58	2424.24	3408.34	1140,38		1526.88	1050 15	3.60	.÷	1145.36	1386.74	1101.20		1175.26	2240.37	2691.70	1410.93		1447.59	1540.64	1869.10	2268.47
Noo No	,	4557371	(NCBI)				•		Q02878	Swiss-	ĵo L	047044°	(Swice-	Prot)			P08112	(Swiss-				Q27967	(Swiss-Prot)				
Cassion	Sprance	human							human			molice	ocnou				human				Commence of the Second	bovine	2 33	-	. ,	• •	
Donal Mood Cago	Profile	5 peaks	match with	BMP-3		10000000000000000000000000000000000000			5 peaks		IIDOSOUTIA LO					Action of the second	4 peaks	match with	· · · · · · · · · · · · · · · · · · ·	4		5 peaks	match with SPP24			1.	
Dana		11	7. ·		3				475				₩. • .				18		, the state of the								

Figure 32 E. Identification of Proteins by Mass Spectrometry of Tryptic Fragments

Mass         Mass Diff- Pas         AAS         % Coversore           Spec         erence         age           Database         -0.11         303-311         63           1175.42         -0.29         400-409         400-409           2084.42         -0.26         312-328         11           2691.91         -0.35         340-362         11           2691.91         -0.37         42-53         11           1447.65         -0.25         113-124         14           1208.40         0.06         48-57         14           1364.59         -0.67         107-118         1364.58           1364.59         -0.07         48-58         102-111           1438.56         -0.02         48-58         102-111           1566.76         0.25         346-357         1651.91           1651.91         -0.10         345-357         109-367			•	· ·						
Spec         Spec         Spec         erence         age           5 peaks         human         Data         Database         -0.11         303-311         63           5 peaks         human         (Swiss-         1101.26         -0.11         303-311         63           TGF-§2         proty         1175.13         1175.42         -0.28         400-409           2240.25         2240.26         -0.26         312-347           2240.25         2240.60         -0.35         340-362           2 peaks         bovine         Q27967         1411.23         1411.60         -0.37         42-53         11           SpP24         proty         proty         1447.40         1447.65         -0.26         113-124           match with mistone H1.x         proty         1221.71         1222.35         -0.64         107-118           1 peaks         proty         12349.86         1360.20         0.05         107-118           1 peaks         proty         1732.23         1732.97         -0.04         48-58           1 peaks         proty         proty         -0.02         107-118         107-118           1 peaks         proty         proty			Species	Acc. No.	Mass	Mass	Mass Diff-	AAs	% Cover-	Collineants
S peaks         human         Database         -0.11         303-311         63           match with artch with histone H1.x         (Swiss-pot)         1101.15         1101.26         -0.28         400-409           2 peaks         2240.25         2240.60         -0.28         312-347           2 peaks         2240.25         2240.60         -0.35         312-328           2 peaks         2091.61         2691.61         2691.91         -0.30         340-362           2 peaks         6 peaks         1417.23         1411.60         -0.37         42-53         11           A peaks         6 peaks         1447.40         1447.65         -0.25         113-124           A peaks         1000         1447.65         -0.25         113-124           A peaks         1221.71         1222.35         -0.64         107-118           A peaks         1000.20         0.05         48-58         144-56           A peaks         11364.57         1364.57         0.07         48-58           A peaks         1000.23         102-111         34-5-37           A peaks         1000.23         102-111         34-5-37           A peaks         140-424         1001.1					Spec	Spec	erence		age	
5 peaks         human         P08112         1101.15         1101.26         -0.11         303-311         63           match with atch with histone H1.x         Prot)         1175.13         1175.42         -0.29         400-409           2084.42         -0.26         312-328         2240.25         2240.60         -0.35         312-328           2 peaks         bovine         Q27967         1411.23         1411.60         -0.37         42-53         11           SPP24         Prot)         1447.40         1447.65         -0.25         113-124         14           Appage         riadch with         Prot)         1447.40         1447.65         -0.25         113-124           Appage         riadch with         Prot)         1447.40         1447.65         -0.25         113-124           Appage         riadch with         Prot)         1447.40         1447.65         -0.25         113-124           Appage         riadch with         Prot)         1427.71         1222.35         -0.07         48-57           Appage         riadch with         Riadch ST         riadch ST         riadch ST         riadch ST           Appage         riadch St         riadch ST         riadch ST	÷ .				Data	Database				
matich with Prof)  TGF- 62  Prof)  1175.13  1175.42  2084.16  2084.16  2084.42  2084.42  2084.42  2084.16  2240.25  2240.25  2240.25  2240.26  2240.27  2240.26  2240	22	5 peaks	human	P08112	1101.15	1101.26	-0.11	303-311	63	
1175.13   1175.42   -0.29   400-409   2084.42   -0.26   312-347   2084.42   -0.26   312-347   2240.25   2240.60   -0.35   312-328   2240.25   2240.60   -0.35   340-382   2240.51   2081.91   -0.30   340-382   111   2240.51   1411.60   -0.37   42-53   111   2691.64   1208.46   1208.40   0.05   48-57   1447.40   1447.45   -0.25   113-124   1349.85   1360.52   -0.67   107-118   1349.85   1360.52   -0.67   107-118   1349.85   1360.52   -0.07   40-58   1360.52   1060.20   0.23   102-111   31   1438.83   1438.58   -0.25   346-357   1566.92   1566.76   0.16   345-357   1566.92   1566.76   0.16   345-357   1566.82   1566.76   0.16   345-357   1566.82   1566.76   1.09   290-318   3407.777   1.09   290-318   140-424   1208.40   140-424		match with		(Swiss-						
2240.25 2240.60 -0.35 312-328 2240.25 2240.60 -0.35 312-328 2240.25 2240.60 -0.35 312-328 2240.25 2240.60 -0.35 312-328  25 peaks		1GF-þ2		<u> </u>	1175 13	1175.42	-0.29	400-409		
2 peaks-match with listone H1.x         bovine G27967         1411.23         1411.60         -0.35         312-328           SPP24 match with listone H1.x         human JC4928         1208.46         1208.40         0.06         48-57         14 match with listone H1.x           Fibeaks match with listone H1.x         human JC4928         1208.46         1208.40         0.06         48-57         14 match with listone H1.x           Fibeaks match with listone H1.x         human JC4928         1208.46         1208.40         0.06         48-57         14 match with listone H1.x           Fibeaks human (NCB)         human H557371         1060.20         0.23         102-111         31           Fibeaks human (NCB)         human H567371         1060.20         0.23         102-111         31           Fibe 92         1566.76         0.16         345-357         1651.91         -0.11         410-424           1566.92         1566.76         1.09         290-318					2084 16	2084.42	-0.26	312-347		
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2 peaks         bovine         Q27967         1411.23         1411.60         -0.37         42-53         11           SPP24         Prof)         1447.40         1447.65         -0.25         113-124           5 peaks         human         JC4928         1208.46         1208.40         0.06         48-57         14           match with match with match with match with lbh-3         human         4557371         1221.71         1222.35         -0.67         107-119           Speaks         human         4557371         1060.43         1060.20         0.23         102-111         31           BMP-3         human         4557371         1060.43         1060.20         0.25         346-357           1566.92         1566.76         0.16         345-357           1566.92         1566.76         0.16         345-357           1651.80         1651.91         -0.11         410-424           3408.86         3407.77         1.09         290-318			:		2691.61	2691.91	-0.30	340-362		
match with SPP24         (Swiss-Frot)         Prot)         1447.40         1447.65         -0.25         113-124           Speaks. human         JC4928         1208.46         1208.40         0.06         48-57         14           match with match with match with match with match with left with match with match with match with match with match with left so 156.92         1364.57         1364.59         -0.074         48-58           Fibeaks         human         4557371         1060.20         0.23         102-111         31           Fibeaks         human         4557371         1060.20         0.25         346-357           1566.76         0.16         345-357         1566.76         0.16         345-357           1566.76         1059. 290-318		2 peaks	bovine	Q27967	1411.23	1411.60	-0.37	42-53	11	
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5-peaks human JC4928 1208.46 1208.40 0.06 48-57 14  match with histone H1.x histone		SFF24		(i)	1447.40	1447.65	-0.25	113-124		
match with histone H1.x	25	5 neaks	hur	JC4928	1208.46	1208.40	0.08	48-57		
1221.71 1222.35 -0.64 107-118 134.57 1364.59 -0.02 48-58 1732.23 1732.97 -0.74 43-57 1060.43 1060.20 0.23 102-111 31 (NCBI) 1438.83 1438.58 0.25 346-357 1566.92 1566.76 0.16 345-357 1651.80 1651.91 -0.11 410-424 3408.86 3407.77 1.09 290-318		match with		(PIR)						
1349,85 1350,52 -0.67 107-119 1364,57 1364,59 -0.02 48-58 1732,23 1732,97 -0.74 43-57 1060,43 1060,20 0.23 102-111 31 (NCB) 1438,83 1438,58 0.25 346-357 1566,92 1566,76 0.16 345-357 1651,80 1651,91 -0.11 410-424 3408,86 3407,77 1.09, 290-318		nistone H1.X			1221.71	1222.35	-0.64	107-118		
human. 4557371 1060.43 1060.20 0.23 102-111 31 (NCBI) 1438.83 1438.58 0.25 346-357 1566.92 1566.76 0.16 345-357 1566.92 1565.76 0.16 345-357 1556.90 1651.91 -0.11 410-424 3408.86 3407.77 1.09, 290-318					1349.85	1350.52	-0.67	107-119		
human 4557371 1060.43 1060,20 0.23 102-111 31 (NCBI) 1060.43 1060,20 0.23 102-111 31 1438.58 0.25 346-357 1566.92 1566.76 0.16 345-357 1651.80 1651.91 -0.11 410-424 3408.86 3407.77 1.09, 290-318					1364.57	1364.59	-0.02			
human. 4557371 1060.43 1060,20 0.23 102-111 31 (NCBI) 1438.83 1438.58 0.25 346-357 1566.92 1566.76 0.16 345-357 1651.80 1651.91 0.011 410-424 3408.86 3407.77 1.09, 290-318					1732.23	1732.97	-0.74	43-57		
(NCB))       1438.83     1438.58       1566.92     1566.76       0.16     345-357       1651.80     1651.91       -0.11     410-424       3408.86     3407.77       1.09     290-318		5 peaks	human	4557371	1060.43	1060.20		102-111	34	% coverage calculation is relative to the mature BMP-
1438.83     1438.58     * 0.25       1566.92     1566.76     0.16       1651.80     1651.91     -0.11       3408.86     3407.77     1.09		match with		SCR SCR						3, 183 AAS (290-472)
1566.76 0.16 1651.91 -0.11 3407.77 1.09		Ì			1438.83	1438.58				.:
1651.91 -0.11 3407.77 1.09					1566.92	1566.76	·			
1.09					1651.80	1651.91	-0.11			,
					3408.86	3407.77				

Figure 32 F. Adentification of Proteins by Mass Spectrometry of Tryptic Fragments

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## FIGURE 33A

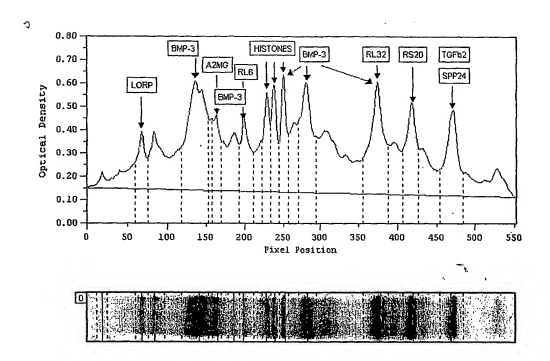


FIGURE 33B

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FIGURE 34: Quantitation of Identified BP proteins

Identified Protein	Percentage of Total Protein
LORP	2
BMP-3	11
BMP-3 and A2-MG	3
RL6 & BMP-3	4
Histone	3
Histone	3
Histone & BMP-3	4
BMP-3	8
RL32 & BMP-3	8
RS2D .	5
SPP24 & TGF-β2	6
Total	58%

nal Application No PCT/US 01/11576

CLASSIFICATION OF SUBJECT MATTER PC 7 A61L27/36 A61L A. CLASS IPC 7 A61L27/50 A61L27/38 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) A61L A61F Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, PAJ C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages 1-11.WO 94 25080 A (MASSACHUSETTS INST γ. 14-59 TECHNOLOGY ; CHILDRENS MEDICAL CENTER (US)) 10 November 1994 (1994-11-10) claims; examples 1-11P,Y WO 00 62832 A (ENDOSPINE LTD) 14-59 26 October 2000 (2000-10-26) claims 1-11, EP 0 411 925 A (UNIV NORTH CAROLINA) γ 14-59 6 February 1991 (1991-02-06) claims 1-11,WO OO 15274 A (GRANT ROY ARTHUR ;OLIVER Α 14-59 ROY FREDERICK (GB); TISSUE SCIENCE LAB LI) 23 March 2000 (2000-03-23) claims; examples Patent family members are listed in annex. Further documents are listed in the continuation of box C. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the A' document defining the general state of the art which is not considered to be of particular relevance 'X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone 'E' earlier document but published on or after the international filing date document which may throw doubts on priority claim(s) or which is clied to establish the publication date of another citation or other special reason (as specified) 'Y' document of particular relevance; the clalmed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the cat. document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed \*&\* document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search 20/08/2001 10 August 2001 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, ESPINOSA, M Fax: (+31-70) 340-3016

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In snal Application No
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#### FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.1

Although claims 10-11, 14-15, 28-30, 37-39 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.

Continuation of Box I.1

Claims Nos.: 12-13

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RULE 23e PCT: HUMAN BODY, ELEMENTS ISOLATED FROM THE HUMAN BODY.

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